# **FINAL**

Remediation by Natural Attenuation Treatability Study for OU-4 (Site FT-1)



Shaw Air Force Base Sumter, South Carolina

**Prepared For** 

Air Force Center for Environmental Excellence Technology Transfer Division Brooks Air Force Base, Texas San Antonio, Texas

and

20 CES/CEVR Shaw Air Force Base Sumter, South Carolina

DTIC QUALITY INSPECTED 4

September 1999

DISTRIBUTION STATEMENT A
Approved for Public Release
Distribution Unlimited

20011030 088

# PARSONS

Parsons Engineering Science, Inc. • A Unit of Parsons Infrastructure & Technology Group Inc.

1700 Broadway, Suite 900 • Denver, Colorado 80290 • (303) 831-8100 • Fax: (303) 831-8208 • www.parsons.com

### 20 September 1999

Mr. Jerry Hansen Technical Program Manager AFCEE/ERT 3207 North Road, Bldg. 532 Brooks AFB, TX 78235-5363

Subject: Submittal of the Final Remediation by Natural Attenuation Treatability Study

for OU-4 (Site FT-1), Shaw AFB, South Carolina (Contract F41624-92-D-

8036)

Dear Mr. Hansen:

Enclosed please find two copies of the September 1999 Final Remediation By Natural Attenuation Treatability Study for OU-4, Shaw AFB, South Carolina. This report was prepared by Parsons Engineering Science, Inc. (Parsons ES) for the Air Force Center for Environmental Excellence Technology Transfer Division (AFCEE/ERT) and the 20 CES/CEV, Shaw AFB, South Carolina.

The intent of the Natural Attenuation Treatability Study was to determine the role of natural attenuation in remediating chlorinated solvent and fuel contamination in groundwater at OU-4. The draft TS was submitted to AFCEE in May 1998. Comments on the draft TS were received from AFCEE as reviewed by Mr. Jon Atkinson of AFCEE (dated 25 February 1999) and Mr. Richard Roller of 20 CES/CEV (dated 18 March 1999). Responses to these comments were prepared by Parsons ES and are attached to this letter. Your response to our comments as submitted to Parsons ES on 18 August 1999 were addressed and these also are attached to this letter.

If you have any questions or require additional information, please call me at (303) 831-8100.

Sincerely,

PARSONS ENGINEERING SCIENCE, INC.

Bruce M. Henry, P.G.

Project Manager

**Enclosures** 

c.c. Mr. Rick Roller – Shaw AFB (two copies)

Mr. Don Kampbell – USEPA NRMRL (two copies)

# PARSONS

Parsons Engineering Science, Inc. • A Unit of Parsons Infrastructure & Technology Group Inc.

1700 Broadway, Suite 900 • Denver, Colorado 80290 • (303) 831-8100 • Fax: (303) 831-8208 • www.parsons.com

### 10 September 1999

Mr. Jerry Hansen Technical Program Manager AFCEE/ERT 3207 North Road, Bldg. 532 Brooks AFB, TX 78235-5363

Subject: Responses to second-round AFCEE Comments on the Draft Remediation by

Natural Attenuation Treatability Study for OU-4 (Site FT-1), Shaw AFB,

South Carolina (Contract F41624-92-D-8036-0006)

Dear Mr. Hansen:

This letter provides responses to second-round comments submitted to Parsons ES by the Air Force Center for Environmental Excellence Technology Transfer Division (AFCEE/ERT) regarding the subject report. On 18 June 1999, Parsons ES submitted to AFCEE/ERT the first response to comments letter regarding the TS report for OU-4 at Shaw AFB. AFCEE/ERT reviewed the 18 June 1999 response to comments letter and submitted additional comments/questions on 18 August 1999. Responses to these second round comments were prepared by Parsons ES and are presented herein.

### **GENERAL COMMENTS**

Comment 1) Can the location of the new reactive barrier wall be added to the site maps?

Parsons ES Response:

Placing the position of the reactive barrier wall into the figures would be incongruent with the data shown on most figures. Groundwater data was collected in May 1997 for this project, whereas the reactive barrier walls were installed in 1998. Therefore, the reactive barrier walls have no significance to plume trends prior to installation. For this reason, Parsons ES recommends that the position of the reactive barrier wall not be placed in every figure. Parsons ES does recommend that a figure of the reactive barrier wall location be placed in Appendix E, and referenced through text changes as a result of comments 2 through 4 below.

- Comment 2) Page ES-3. SVE may not be required if the wall works. Recommend that SVE be implemented only if the wall does not adequately reduce the discharge to the creek. There is no guarantee that SVE can clean up the source to a point where we would see a reduction in LTM etc.
- Parsons ES Response: Agreed. Parsons ES concurs that RNA, the permeable reactive barrier wall, and LTM are sufficient to prevent CAH discharge to Long Branch Creek at rates above set regulatory standards. Since natural source weathering will be the only mechanism that will reduce source mass, the duration of an LTM program may continue for decades, although lowered sampling frequencies may be achieved during this period. Section 6 was modified to recommend RNA, LTM, institutional controls, and the continued operation of the permeable reactive barrier wall (modified Alternative 1) versus the implementation of an SVE system (Alternative 2). This change was applied to Page ES-3 to coincide with text revisions in Section 6. Attached pages of the report show page ES-3 and Section 6 text changes in bold.
- Comment 3) Page 6-17. Discussion on funnel and gate may not be relevant if the current existing wall covers the total plume flow. Since the wall is already there it must be considered in the final recommendation. The report can indicate that the analysis of alternatives did not look at a wall, but that the wall was installed under a separate effort.
- <u>Parsons ES Response:</u> Reference to the funnel gate was deleted from page 6-17.
- Comment 4) Page 6-42. SVE may be a consideration down the road if the wall fails etc. Current course of action should be to continue to monitor the site for NA, reactive wall performance, and source area weathering. That may turn out to be all that is needed.
- <u>Parsons ES Response:</u> Agreed. The use of SVE as a contingency action was incorporated into the report as per the response to comment 2.
- Comment 5) Page 8-3. Words indicate a "planned groundwater extraction and treatment system to be installed at the site". Do not indicate that the system is planned. Indicate that a model was used to simulate a P&T only as another alternative.
- Parsons ES Response: Agreed. The referenced sentence was changed to "A third groundwater model was used to simulate the potential effects-of the planned of a groundwater extraction and treatment system to be installed at the site."

Mr. Jerry Hansen 10 September 1999 Page 3

If you have any questions, have additional comments, or require additional information, please call me at (303) 831-8100.

Sincerely,

PARSONS ENGINEERING SCIENCE, INC.

Bruce M. Henry, P.G. Project Manager

cc: Mr. R. Todd Herrington, Parsons ES File 722450.37000

# FINAL TREATABILITY STUDY IN SUPPORT OF REMEDIATION BY NATURAL ATTENUATION FOR GROUNDWATER AT OU-4 (SITE FT-1)

at

SHAW AIR FORCE BASE SOUTH CAROLINA

**SEPTEMBER 1999** 

### Prepared for:

AIR FORCE CENTER FOR ENVIRONMENTAL EXCELLENCE
TECHNOLOGY TRANSFER DIVISION
BROOKS AIR FORCE BASE
SAN ANTONIO, TEXAS

and

20 CES/CEVR SHAW AIR FORCE BASE SOUTH CAROLINA

### Prepared by:

PARSONS ENGINEERING SCIENCE, INC. 1700 BROADWAY, SUITE 900 DENVER, COLORADO 80290

## TABLE OF CONTENTS

			Page
ACRC	NYMS	AND ABBREVIATIONS	ix
EXEC	UTIVE	SUMMARY	ES- 1
SECT	ION 1 -	INTRODUCTION	
1.1 1.2 1.3 1.4 1.5	Report Installa OU-4	Organizationtion Description and Hist Background	
SECT	ION 2 -	SITE CHARACTERIZA	TION ACTIVITIES 2-1
2.1	Cone 1 2.1.1	Monitoring Point Placem Stratigraphy with CPT	ent and Investigation of Site
	2.1.2	Investigation of Residual	and Mobile Hydrocarbons2-2
2.2	Geopr	be Field Activities	2-3
2.3	Groun	water Monitoring Point (	Completion
	2.3.1		ıction
			Activities
			contamination Procedures
			int Materials Decontamination
	2.3.2		tion
	2.3.3		pment2-7
2.4			
2	2.4.1	Groundwater Sampling I	ocations 2-8
			nt Cleaning
	2.4.3		rocedures
			Location2-9
		2.4.3.2 Water Level as	nd Total Depth Measurements 2-9
		2.4.3.3 Monitoring Wo	ell/Point Purging2-10
			ion2-10
	244	Onsite Chemical Paramet	er Measurement 2-10

		J.	Page
	2.4.5 Sample	le Handling	2-11
2.5	Surface Water	er Sampling	2-11
2.6		npling	
2.7		pling	
2.8		ing	
	1		
SECT	ION 3 - PHYS	SICAL CHARACTERISTICS OF THE STUDY AREA	3-1
3.1	Topography,	Surface Hydrology, and Climate	3-1
3.2		ology and Hydrogeology	
3.3	Site Geology	and Hydrogeology	3-5
	3.3.1 Lithole	logy and Stratigraphic Relationships	3-5
	3.3.2 Groun	ndwater Hydraulics	3-8
	3.3.2.		3-8
	3.3.2.	.2 Hydraulic Conductivity	3-11
		.3 Effective Porosity	
	3.3.2.	.4 Advective Groundwater Velocity	3-11
3.4	Groundwater	Use	3-12
SECT	ION 4 - CONT	TAMINANT DISTRIBUTION AND EVIDENCE FOR	
	BIOD	DEGRADATION	. 4-1
4.1	Contaminant !	Sources and Soil Quality	4-1
7.1	4.1.1 Mobile	le LANPL Contamination	4-1
		Contamination	
4.2	Overview of 1	Hydrocarbon Biodegradation	4-6
	4.2.1 Review	w of Biodegradation Processes	4-6
		gradation of BTEX	
	4.2.3 Biodes	gradation of Chlorinated Solvents	4-8
	4.2.3.		
		Dehalogenation)	. 4-9
	4.2.3.		
	4.2.3.		4-11
	4.2.3. 4.2.3. 4.2.3.	.3 Cometabolism	4-12
	4.2.3.	.3 Cometabolism	4-12 4-12 4-12
	4.2.3.	.3 Cometabolism	4-12 4-12 4-12 4-14
	4.2.3.	.3 Cometabolism	4-12 4-12 4-12 4-14 4-14
	4.2.3.	.3 Cometabolism	4-12 4-12 4-12 4-14 4-14

				Page
4.3	Distril		Hydrocarbons and Daughter Products	
	4.3.1	Fuel Hyd	lrocarbons	4-16
	4.3.2	Chlorina	ted Ethenes	4-18
	4.3.3	Chlorina	ted Ethanes	4-24
	4.3.4	Chlorobe	enzenes	4-24
	4.3.5	Other Ch	llorinated Compounds	4-28
	4.3.6	Chloride		4-28
4.4	Additi	onal Evide	ence of Biodegradation	4-31
	4.4.1	Electron	Donors, Native Electron Acceptors, and Byproducts	4-31
		4.4.1.1	Organic Carbon in Groundwater	4-31
		4.4.1.2	Inorganic Chemistry	
			4.4.1.2.1 Dissolved Oxygen	
			4.4.1.2.2 Nitrate/Nitrite	
			4.4.1.2.3 Sulfate	
-			4.4.1.2.4 Ferrous Iron	
			4.4.1.2.5 Methane	
			4.4.1.2.6 Ammonia/Ammonium	
		4.4.1.3	Ethane/Ethene in Groundwater	
	4.4.2		al Geochemical Indicators	
		4.4.2.1	Oxidation/Reduction Potential and Dissolved	
			Hydrogen as Indicators of Redox Processes	
		4.4.2.2	Alkalinity and Carbon Dioxide Evolution	
		4.4.2.4	pH	
		4.4.2.5	Temperature	
4.5			of Biodegradation Rates	
4.6	Discus	ssion		4-46
aram	TON E	CDOLIN	DULL TED EL OUL AND CONTANINANT	
SECT	ION 3		DWATER FLOW AND CONTAMINANT	5 1
		IKAN5	PORT MODEL	3-1
5.1	Gener	al Overvie	ew and Model Description	5_1
5.2	Conce	ar Overvie	del Design and Assumptions	5-2
5.3	Initial	Model Se	tup	5-3
3.3			sign	
			vater Flow Model	
	9.9.2	5.3.2.1		
		5.3.2.2		
		5.3.2.3	Aquifer Properties	
	5.3.3		nant Transport Model	

			Page
		5.3.3.1 Source	5-7
		5.3.3.2 Dispersivity	5-8
		5.3.3.3 Retardation	
		5.3.3.4 Biodegradation	
5.4	Model	Calibration	5-9
	5.4.1		5-9
	5.4.2	Plume Calibration	
5.5		vity Analysis	5-14
	5.5.1	Sensitivity to Variations in Hydraulic Conductivity	5-15
	5.5.2	Sensitivity to Variations in Dispersivity	5-15
	5.5.3	Sensitivity to Variations in the Coefficient of Retardation	5-16
	5.5.4	Sensitivity to Variations in the Anaerobic Decay Rate Constant	
	5.5.6	Summary of Sensitivity Analysis Results	
SECT	ION 6 -	COMPARATIVE ANALYSIS OF REMEDIAL	
		ALTERNATIVES	6-1
6.1	Remed	lial Alternative Evaluation Criteria	6-1
	6.1.1	Long-Term Effectiveness and Permanence	6-2
	6.1.2	Implementability	6-2
	6.1.3		6-2
6.2		s Influencing Alternatives Development	
	6.2.1	Program Objectives	6-3
	6.2.2	Contaminant Properties	6-3
	6.2.3	Site-Specific Conditions	6-4
		6.2.3.1 Groundwater and Soil Characteristics	
		6.2.3.2 Potential Exposure Pathways	6-6
		6.2.3.3 Remediation Goals for Shallow Groundwater	
	6.2.4		6-8
6.3		Description of Remedial Alternatives	6-14
	6.3.1	Alternative 1RNA and Institutional Controls with Long-Term	
		Groundwater and Surface Water Monitoring	6-14
	6.3.2	Alternative 2—SVE of the 1,1,1-TCA Source, RNA, and	
		Institutional Controls with Long-Term Groundwater and Surface	
		Water Monitoring	6-15
6.4		ation of Alternatives	6-16
	6.4.1		
		Groundwater and Surface Water Monitoring	
		6.4.1.1 Effectiveness	6-16

				Page
		6.4.1.2	Technical and Administrative Implementability	
		6.4.1.3	Cost	6-21
	6.4.2	Alternativ	ve 2 - SVE of the 1,1,1-TCA Source, RNA, and	
		Institution	nal Controls with Long-Term Groundwater and Surface	
			onitoring	6-21
		6.4.2.1	Effectiveness	
		6.4.2.2	Technical and Administrative Implementability	6-24
			Cost	
6.5			imp-and-Treat As An Interim Measure To Monitored	
			ion	
6.6	Recon	mended R	Remedial Approach	6-30
an am		T ONG T	EDICIONITODING DI AN	7.1
SECT	ION 7 -	LONG-1	ERM MONITORING PLAN	/-1
7.1 -	Overv	iow		7-1
7.2			works	
1.2	7 2 1	Groundw	vater Monitoring Network and Sampling Frequency	7-2
			ells	
7.3	Surfac	e Water M	Ionitoring Network and Sampling Frequency	7-2
7.4			col	
7.5			lan Review	
SECT	ON 8 -	CONCLU	USIONS AND RECOMMENDATIONS	8-1
anami		DEFEDE	PICEG	0.1
SECT	ION 9 -	REFERE	INCES	9-1
APPE	NDIX A	A - Boreh	ole logs, CPT/LIF results, Monitoring Well/Point	
		Samplin	ng Forms, Aquifer Test Results, and Survey Data.	
V DDE.	NDIY I	B – Analyt	rical Data	
		•		
APPE	NDIX (	C - Model	ing Information	
APPE	NDIX I	D - Model	Input Files	
APPE	NDIX I	E – Cost C	Calculations	

## LIST OF TABLES

Title	Page
History of Past Remedial Activities at OU-4	. 1-6
Summary of Well Installation Details	. 2-4
Analytical Protocol used for Groundwater, Surface Water, Sediment,	
and Mobile LNAPL Samples	. 2-8
Summary of Groundwater Elevation Data May 1997	. 3-9
Summary of Detected Volatile and Diesel-Range Organic Compounds	
	. 4-4
Groundwater Quality Data Summary for BTEX, TMBs, and Fuel	
Carbon	4-17
	4-21
Groundwater Chemical Parameter	
Range of Hydrogen Concentrations for a Given Terminal Electron-	
Accepting Process	4-42
Summary of Chlorinated Solvent and BTEX Decay Rates	4-45
Analytical Parameters And Weighting For Preliminary Screening OU-4	
RNA TS Shaw AFB, South Carolina	4-49
Interpretation Of Points Awarded During Natural Attenuation	
Screening	
Common Designations For Several Important Boundary Conditions	. 5-5
Calculation of Retardation Coefficients (Foc=0.00024)	
	. 6-8
Initial Technical Implementability Screening of Technologies and	
Estimated Alternative 2 Costs	
Long-Term Groundwater Monitoring Analytical Protocol	. 7-5
	History of Past Remedial Activities at OU-4 Summary of Well Installation Details Analytical Protocol used for Groundwater, Surface Water, Sediment, and Mobile LNAPL Samples Summary of Groundwater Elevation Data May 1997 Summary of Detected Volatile and Diesel-Range Organic Compounds in OU-4 Soil Samples Collected in February/March 1993 Groundwater Quality Data Summary for BTEX, TMBs, and Fuel Carbon Groundwater and Surface Water Quality Data summary for Chlorinated Ethenes, Ethanes, and Chloroform Groundwater Quality Data Summary for Chlorinated Benzenes Groundwater Electron Acceptors/Byproducts Groundwater Chemical Parameter Range of Hydrogen Concentrations for a Given Terminal Electron-Accepting Process Summary of Chlorinated Solvent and BTEX Decay Rates Analytical Parameters And Weighting For Preliminary Screening OU-4 RNA TS Shaw AFB, South Carolina Interpretation Of Points Awarded During Natural Attenuation Screening. Common Designations For Several Important Boundary Conditions Calculation of Retardation Coefficients (Foc=0.00024) Sensitivity Analysis Results. Remedial Action Objectives For Groundwater And Surface Water Initial Technical Implementability Screening of Technologies and Process Options for Groundwater Remediation Estimated Alternative 1 Costs

### LIST OF FIGURES

No.	Title	Page
1.1	Base Location	1-4
1.2	Site Location	1-9
1.3	Former Fire Training Area 1, OU-4 Layout	1-10
2.1	Sampling Locations	2-5
3.1	Surface Water Drainage	3-2
3.2	Hydrogeologic Cross-Section for Shaw AFB	3-4
3.3	Hydrogeologic Cross-Section A-A'	3-6
3.4	Hydrogeologic Cross-Section B-B'	3-7
3.5	Shallow Groundwater Elevations May 1997	3-10
4.1	Extent of Mobile LNAPL and VOCs in Soil	4-3
4.2	Anaerobic Reductive Dehalogenation	4-10
4.3	Aerobic Degradation	4-13
4.4	Total BETEX and Fuel Carbon Measured a Shallow Monitoring	
	Wells/Points May 1997	4-19
4.5	BTEX Compounds Measured at Shallow Monitoring Wells/Points May	
	1997	4-19
4.6	Chlorinated ethene Contamination at Shallow Monitoring Wells/Points	
		4-22
4.7	Chlorinated Ethanes, 1,1-DCE, and Chloroform Measured at Shallow	
		4-25
4.8	Chlorinated Benzene Contamination Measured at Shallow Monitoring	
		4-27
4.9	Dissolved Chlorine Measured at Shallow Monitoring Well/Point May	4.00
4 10		4-30
4.10	Dissolved TOC Concentrations Measured at Shallow Monitoirng	4.22
4.11	Wells/Points May 1997  Electron Acceptor Trends Measured at Shallow Monitoirng	4-33
4.11	Wells/Points May 1997	1 25
4.12	Dissolved Metabolic Byproduct Concentrations Measured at Shallow	4-33
7.12	Monitoring Wells/Points May 1997	1_37
4.13	Ethane/Ethene Concentrations Measured at Shallow Monitoring	4-37
7.13	Wells/Points May 1997	4-40
4.14	Oxidation Reduction Potentials at Shallow Monitoring Wells/Points	
	May 1997	4-41
5.1	Model Grid	

## LIST OF FIGURES (Continued)

No.	Title	Page
5.2	Total CAH Plume	5-7
5.3	Calibrated Groundwater Elevations	5-11
5.4	Calibrated Total CAH Plume	5-13
6.1	Simulation of Total CAH Concentrations using Model OU4MODA	
	Natural Attenuation Only	6-18
6.2	Simulation of 1,1,1-TCA Concentrations using Model OU4MODA	
	Natural Attenuation Only	6-18
6.3	Plot of 1,1,1-TCA Concentrations Versus Time for Model OU4MODA	6-20
6.4	Simulation of Total CAH Concentrations using Model OU5MODA	
	Natural Attenuation with Soil Vapor Extraction in the Source Area	6-23
6.5	Simulation of 1,1,1-TCA Concentrations using Model OU5MODA	
	Natural Attenuation with Soil Vapor Extraction in the Source Area	6-25
6.6	Plot of 1,1,1-TCA Concentrations Versus Time for Model OU4MODA	6-25
6.7	Simulation of 1,1,1-TCA Concentrations using Model OU4MODC-	
	Natural Attenuation with 2 Years Groundwater Extraction at 1,1,1-	
	TCA Source	
7.1	Proposed Long-Term Monitoring Locations	7-3

### ACRONYMS AND ABBREVIATIONS

μg/kg micrograms per kilogram

 $\Delta G^{\circ}r$  Gibbs free energy of reaction

 $\begin{array}{ll} \mu g/L & \text{micrograms per liter} \\ 1,2\text{-DCE} & 1,2\text{-Dichloroethylene} \\ ACC & \text{Air Combat Command} \end{array}$ 

AFB Air Force Base

AFCEE United States Air Force Center for Environmental Excellence

CAH chlorinated aliphatic hydrocarbon

CB chlorobenzene
CPT Cone Penetrometer
DCA dichloroethane
DCB dichlorobenzene
DRO diesel range organics
ES Engineering Science

FS/CMS Feasibility Study/Corrective Measure Study

ft/day feet per day ft/ft foot per foot

HDPE high density polyethylene IT International Technology

IWTP industrial wastewater treatment plant

LAW Law Environmental

LIF laser induced fluorescence LNAPL light nonaqueous phase liquid

LTM long-term monitoring

MCL maximum contaminant level mg/kg milligrams per kilogram

NRMRL National Risk Management Research Laboratory

ORD Office of Research and Development

OSWER · Office of Solid Waste and Emergency Response

OU-4 Operable Unit 4
PCA tetrachloroethane
PCE tetrachloroethene

POL petroleum, oils, and lubricants

PVC polyvinyl chloride QC quality control

RAP Remedial Action Plan

RI/FS Remedial Investigation/Feasibility Study RNA remediation by natural attenuation Rust E&I Rust Environment and Infrastucture

SCDHEC South Carolina Department of Health & Environmental Control

SS stainless steel
TCA trichloroethane
TCB trichlorobenzene
TCE trichloroethene

TeCB	tetrachlorobenzene
TMB	trimethylbenzene
TS	Treatability Study
USACE	United States Army Corps of Engineers
USEPA	United States Environmental Protection Agency
VC	vinyl chloride

### **EXECUTIVE SUMMARY**

This report presents the results of a treatability study (TS) performed by Parsons Engineering Science, Inc. (Parsons ES) at Operable Unit 4 (OU-4) at Shaw Air Force Base, South Carolina, to evaluate remediation by natural attenuation (RNA) of dissolved chlorinated aliphatic hydrocarbons (CAHs) and fuel hydrocarbons in groundwater. The TS focused on the fate and transport of dissolved CAHs, particularly 1,1,1-trichloroethane (1,1,1-TCA), which is present at high concentrations at the site. Petroleum hydrocarbons [including benzene, toluene, ethylbenzene, and xylenes (BTEX)] also are present in site groundwater and were considered in this TS. The site history and the results of the soil and groundwater investigations conducted previously also are summarized in this report.

Comparison of CAH, BTEX, electron acceptor, and biodegradation byproduct isopleth maps for OU-4 provides strong qualitative evidence of biodegradation of dissolved CAHs. These geochemical data strongly suggest that reductive dehalogenation of dissolved CAHs is occurring at the site as microbes utilize natural organic carbon, BTEX, and other fuel hydrocarbons as substrates. Patterns observed in the distribution of CAHs, daughter products of CAHs, electron acceptors, and biodegradation byproducts also suggest that dehydrohalogenation may be acting to reduce the concentration of CAHs dissolved in site groundwater. Field-scale, firstorder decay rates computed using data from OU-4 include a total BTEX biodegradation rate of 0.014 day<sup>-1</sup>, a total chlorinated ethene decay rate of 0.012 day<sup>-1</sup>, a total chlorinated ethane decay rate of 0.013 day<sup>-1</sup>, and a dichlorobenzene decay rate of 0.011 The equivalent half-lives are 50 days, 56 days, 53 days, and 61 days, respectively. The computed rate of reductive dechlorination for total chlorinated ethene and ethane contamination is 0.0026 day<sup>-1</sup> (half-life of 266 days) and 0.0025 day<sup>-1</sup> (halflife of 213 days), respectively.

An important component of this study was an assessment of the potential for contamination in groundwater to migrate from the source areas to potential receptor exposure points (i.e., stretches of creeks or ponds downgradient or downstream from the source). The finite-difference models MODFLOW and MT3D96® were used in conjunction with site-specific geologic, hydrologic, and laboratory analytical data to simulate the migration and biodegradation of CAHs dissolved in groundwater. Model parameters that were not measured at the site were estimated using reasonable literature values.

The results of this demonstration indicate that RNA of dissolved CAHs and BTEX contamination is occurring at OU-4. Conservative modeling suggests that under current conditions, the CAH plume is at steady-state, and dissolved CAHs will continue to migrate from the source area to the edge of Long Branch Creek that borders OU-4 to the east. The estimated rates of biodegradation combined with the effects of sorption, dispersion, and dilution result in a two-order-of-magnitude reduction in dissolved CAH concentrations between the source area and Long-Branch Creek. CAH concentrations discharging to Long Branch Creek are significantly lower than source area concentrations, and appear to have minimal impact on surface water quality. Maximum groundwater CAH concentrations coincide with the presence of observed light non-aqueous phase liquid (LNAPL) in monitoring points installed at OU-4. Therefore,

model results indicate the importance of source removal to assist natural attenuation processes for site remediation.

Future site activities are not anticipated to change, and the risk to any Base personnel would be minimal, provided institutional controls for soil and groundwater content are maintained. Institutional controls such, as restrictions on shallow groundwater and surface water use at the site would prevent completion of receptor exposure pathways until RNA is complete.

A permeable reactive barrier wall (iron filings trench) was installed in November 1998 to mitigate the migration of contaminants to Long Branch Creek to the east of OU-4. The permeable reactive barrier wall is a passive technology intended to degrade dissolved contaminants migrating from the site. The primary focus of this TS is to evaluate the use of RNA, both alone and in combination with engineered remedial actions, to minimize LTM periods and remediation costs. Based on the evaluation of site contamination at OU-4, the Air Force Center for Environmental Excellence (AFCEE) recommends implementation of RNA, long-term monitoring (LTM), institutional controls, and continued operation of the permeable reactive barrier wall.

To verify model predictions, and to ensure that the selected technologies are meeting objectives, the Air Force recommends using periodic sampling of 11 LTM wells, 1 sentry well, and 6 surface water locations to monitor the long-term migration and degradation of the dissolved CAH plume. In addition to analyses used to verify the effectiveness of RNA, the groundwater samples should be analyzed for CAH and BTEX compounds by US Environmental Protection Agency (USEPA) Method SW8260b or SW8021b. If data collected under the LTM program indicate that the selected remedial alternative is not sufficient to reduce CAH concentrations, specifically 1,1,1-TCA concentrations, at downgradient well locations to levels considered protective of human health and the environment, additional corrective actions may be required to remediate groundwater at the site.

### **SECTION 1**

### INTRODUCTION

This report presents the results of a treatability study (TS) conducted to evaluate the potential effectiveness of remediation by natural attenuation (RNA) for groundwater contaminated with several types of chlorinated aliphatic hydrocarbons (CAHs) and petroleum hydrocarbons at the Former Fire Training Area 1, Operable Unit 4 (OU-4) at Shaw Air Force Base (AFB) in Sumter, South Carolina. As used in this report, RNA refers to a management strategy that relies on natural attenuation mechanisms to remediate contaminants dissolved in groundwater and to control receptor exposure risks associated with contaminants in the subsurface. The United States Environmental Protection Agency (USEPA) Offices of Research and Development (ORD) and Solid Waste and Emergency Response (OSWER) define natural attenuation as (Wilson, 1996):

The naturally occurring processes in soil and groundwater environments that act without human intervention to reduce the mass, toxicity, mobility, volume, or concentration of contaminants in those media. These in-situ processes include biodegradation, dispersion, dilution, adsorption, volatilization, and chemical or biological stabilization or destruction of contaminants.

As suggested by this definition, mechanisms for natural attenuation of CAHs include advection, dispersion, dilution from recharge, sorption, volatilization, abiotic chemical transformation, and biodegradation. Of these processes, biodegradation is the most common mechanism working to transform contaminants into innocuous byproducts. During natural biodegradation, indigenous microorganisms work to bring about a reduction in the total mass of contamination in the subsurface without artificial intervention (e.g., addition of nutrients). Patterns and rates of natural attenuation can vary markedly from site to site depending on governing physical and chemical processes.

### 1.1 SCOPE AND OBJECTIVES

Parsons ES was retained by the United States Air Force Center for Environmental Excellence (AFCEE) Technology Transfer Division to conduct site characterization and groundwater modeling to evaluate the scientific defensibility and technical feasibility of RNA with long-term monitoring (LTM) as a remedial option for fuel- and industrial-solvent- contaminated groundwater at OU-4. Site characterization activities conducted at OU-4 as part of this study consisted of numerous tasks that were required to fulfill the project objective. These tasks included:

- Reviewing existing hydrogeologic and soil and groundwater quality data for the site;
- Conducting supplemental site characterization activities to more thoroughly characterize the nature and extent of groundwater contamination;
- Collecting geochemical data in support of RNA;
- Developing a conceptual hydrogeologic model of the shallow saturated zone, including the current distribution of contaminants;
- Evaluating site-specific data to determine whether natural contaminant attenuation and destruction are occurring in groundwater at the site;
- Designing and executing a groundwater flow and contaminant fate and transport model for site hydrogeologic conditions;
- Simulating the fate and transport of the total chlorinated solvent plume (e.g., chlorinated ethenes, ethanes and benzenes) in groundwater under the influence of advection, dispersion, adsorption, and biodegradation using the calibrated model;
- Evaluating a range of model input parameters to determine the sensitivity of the model to those parameters and to consider several contaminant fate and transport scenarios;
- Determining if natural processes are minimizing plume expansion of dissolved contaminants so that groundwater and surface water quality standards can be met downgradient;
- Conducting an exposure pathways analysis for potential current and future receptors; and
- Using the results of modeling to determine the adequacy of RNA to protect human health and the ecosystem and to formulate a LTM plan.

A two-phase site characterization effort was conducted to collect the data necessary to support RNA at OU-4. Phase I site characterization activities were completed on April 3-4, 1997, and involved the exploration of the subsurface with cone penetrometer testing (CPT) and laser-induced fluorescence (LIF) equipment, and placement of five groundwater monitoring points (MPA through MPE). Phase II site characterization was completed from May 14 through 18, 1997, and involved installation of six additional monitoring points (TMP 1 through TMP 6); reinstallation of monitoring point MPE with a Geoprobe®; groundwater sample collection from site monitoring wells and points; surface water and sediment sample collection from site surface water bodies; analysis of groundwater, surface water and sediment samples; collection and analysis of mobile light nonaqueous-phase liquid (LNAPL, or free product) from site monitoring points; aquifer testing; and static groundwater level measurement. Field investigation methods are described in the TS Work Plan (Parsons ES, 1997).

Site-specific data collected during the May 1997 field investigation were used to develop a groundwater flow and solute fate and transport model for the site and to conduct a preliminary exposure pathways analysis. The model was used to predict the future extent and concentrations of dissolved CAH and chlorinated aromatic hydrocarbon plumes by simulating the combined effects of advection, dispersion, sorption, and biodegradation. Results of the model were used to predict future discharge to surface water, to assess the potential for completion of other exposure pathways involving groundwater, and to identify whether natural attenuation with LTM is an appropriate and defensible remedial option for contaminated groundwater.

This report is intended to be used by the Base, the US Army Corps of Engineers (USACE), the USEPA, the South Carolina Department of Health and Environmental Control (SCDHEC), and other agencies responsible for establishing and approving a site closure plan. Specifically, the results presented herein can be used to justify the future role of RNA with LTM in site remediation, either as the sole remediation alternative or in conjunction with engineered remedial alternatives proposed in the feasibility study/corrective measures study (FS/CMS) for the site [Rust Environment & Infrastructure (Rust E&I), 1996].

### 1.2 REPORT ORGANIZATION

This TS contains nine sections, including this introduction, and four appendices. Section 2 summarizes site characterization activities. Section 3 summarizes the physical characteristics of the study area. Section 4 describes the nature and extent of soil, groundwater, surface water, and sediment contamination, and the geochemistry of soil and groundwater at the site. Section 5 describes the design of the conceptual hydrogeologic model and numerical groundwater flow and contaminant fate and transport model for the site; lists model assumptions and input parameters; and describes sensitivity analyses, model output, and the results of the modeling. Section 6 presents a comparative analysis of remedial alternatives, and Section 7 presents the LTM plan for the site. Section 8 presents the conclusions of this work and provides recommendations for further work at the site. Section 9 lists the references used to develop this document. Appendix A contains borehole logs, CPT/LIF results. monitoring well construction diagrams, monitoring well/point sampling forms, aquifer test results, and survey data. Appendix B presents soil, sediment, groundwater, and surface water analytical results collected as part of this TS. Appendix C contains model input parameters, model output and supporting figures, and calculations related to model calibration. Appendix D contains model input files. Appendix E contains cost calculations pertinent to remedial alternatives selection. Appendix F contains a figure of the iron filings wall installed in November 1998 at Site S-1. Appendix G contains review comments and responses for this report.

### 1.3 INSTALLATION DESCRIPTION AND HISTORY

Shaw AFB is located near the center of South Carolina in northwest Sumter County, approximately 37 miles east of Columbia and 7 miles west of the city of Sumter along US Highway 76 (Figure 1.1). The Base encompasses 3,326 acres of land set in a mostly rural and agricultural area; however, residential and commercial developments are present in the western and southeastern parts of the Base (Rust E&I, 1995).

Established as an Air Force flying school in 1941, the Base was under the Tactical Air Command. Numerous aircraft changes have since occurred at the Base, but the mission has always been to provide tactical fighter forces. To support this mission, quantities of petroleum, oils, and lubricants (POL); solvents; and coatings were used, resulting in waste generation. The host organization for the Base is the 20th Fighter Wing, which has three F-16 Squadrons. In addition, several tenant units are located at the Base. The Base and its fighter wing are now under the Air Combat Command (ACC).

### 1.4 OU-4 BACKGROUND

OU-4 encompasses the area impacted by activities at Former Fire Training Area 1. The site consists of approximately 14 acres located between Patrol Road on the west and Long Branch Creek (a surface water drainage along the east-northeastern Base property boundary) on the east (Figures 1.2 and 1.3). The remnants of a berm provide evidence of a former burn pit; however, interpretation of historical aerial photographs suggests that the fire pit was moved periodically during the operation of the facility (Rust E&I, 1995). The site features sparse vegetation except in the heavily wooded perimeter and on the slopes near Long Branch Creek.

Reportedly, the former fire training area was used for weekly fire training exercises from 1941 until 1969. A variety of combustible wastes were burned during the training exercises, including jet fuel, waste oils, hydraulic fluids, spent solvents, contaminated fuels, and napalm. Waste materials were hauled to the site in drums and poured into unlined burn pits for ignition. Extinguishing agents reportedly used during training exercises included water, carbon dioxide, protein foam, and aqueous-film-forming foam. Some of the drums used to store and transport the combustibles may have been buried at the site (Parsons ES, 1994; Rust E&I, 1995).

The fire training area was first identified as having a "high potential for environmental contamination" in a 1983 Phase I assessment [Engineering-Science, Inc. (ES). 1983). Since the initial Phase I assessment, numerous remedial activities have been performed at the site; these activities are summarized in Table 1.1 and discussed below. A Phase II, Stage I investigation was performed by Research Triangle Institute (RTI, 1986) in January 1986 and a Stage II investigation was performed by Law Environmental, Inc. (Law, 1989) between July 1986 and February 1987. A remedial action plan (RAP) was prepared by Law (1989) to identify, evaluate, and recommend remedial action alternatives that would address the contamination at the site. At that time, a groundwater monitoring program was selected as the alternative capable of providing the appropriate level of protection to the environment and public health at the most reasonable cost. After collection of two years of quarterly groundwater monitoring data (1988 - 1990), it was decided that long-term monitoring did not adequately address site remediation and that additional investigations were warranted. Since reopening the investigation, Law (1991) prepared a draft remedial investigation (RI)/FS, and Rust E&I (1995) conducted supplemental RI activities and issued the final RI report.

# HISTORY OF PAST REMEDIAL ACTIVITIES AT OU-4

OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

Activity	Report/Reference	Results/Recommendations
Identify contaminated Sites Shaw AFB	Installation Restoration Program (IRP), Phase I (Engineering Science, 1983)	OU-4 was identified as having "high potential for environmental contamination" during Phase 1 study. Site recommended for Phase II investigation to provide more specific contaminant information.
Verify the presence of site contamination at OU-4	IRP, Phase II, Stage I (RTI, 1986)	Four wells installed at OU-4 (MW-105, -106, -107, and -111, see Figure 1.3) to measure potential downgradient and background contaminant concentrations. Analytical data verified that the shallow aquifer had been impacted by oil and grease, total organic halogens, and various volatile organic compounds (VOCs).
Further delineate soil and groundwater contamination at OU-4	IRP, Phase II, Stage II; July 1986 to February 1987 (Law, 1989)	On the basis of Phase II, Stage 1 information, Law Environmental installed five shallow monitoring wells (MW-112, -113, -113A, -114, and -115), three deep (Black Creek Aquifer) wells (MW-105D, -106D, and -112D), a pump test well (TW-101); and three observation wells (PZ-101, -102, and -103); two surface water and sediment locations (SW/SD-11 and -12); and one soil boring within the fire pit berm (SB-111). Soil data indicated that total recoverable petroleum hydrocarbon (TRPH) and lead contamination occurred primarily in the vadose zone beneath the fire training pit. A pump test was performed at TW-101 using observation wells PZ-101, -102, and -103.
Perform quarterly groundwater and surface water monitoring at OU-	Phase II sampling; 1988 through 1990 (Law, 1989 and 1991).	Shallow aquifer determined to be contaminated with benzene, toluene, ethylbenzene, and xylenes (BTEX), chlorinated solvents, and metals. Two surface water stations (SW-1-3 and SW1) were included in the quarterly monitoring program. BTEX compounds in groundwater and saturated sediments indicated that contaminants had migrated from source area to Long Branch Creek. No VOCs other than methylene chloride were detected in surface water.

# TABLE 1.1 (Continued) HISTORY OF PAST REMEDIAL ACTIVITIES AT OU-4 OUTA BNA TS

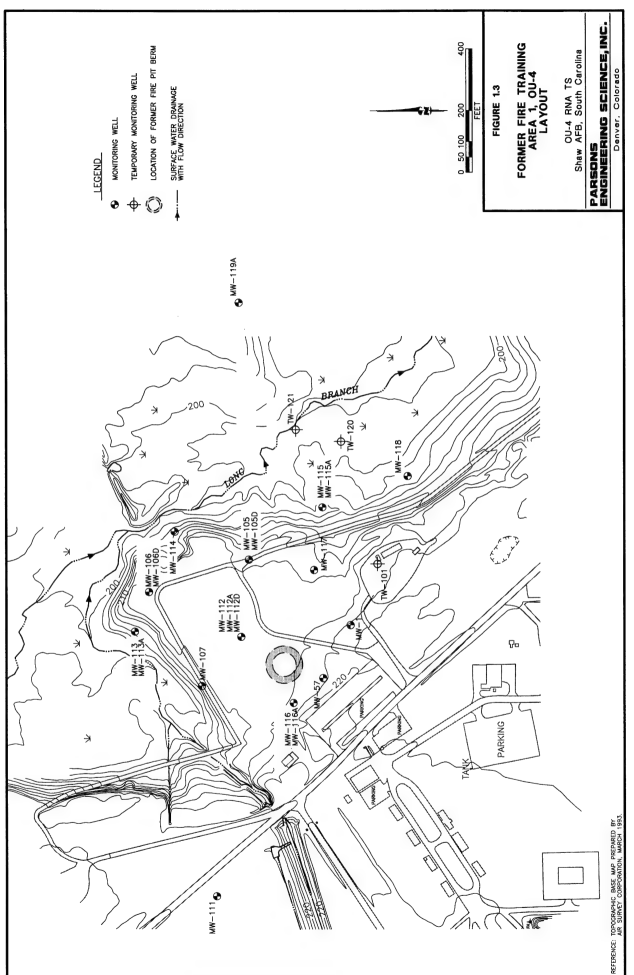
OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

Activity	Report/Reference	Results/Recommendations
Identify, evaluate, and recommend remedial actions	Remedial Action Plan (Law, 1989)	On the basis of Phase II site data, groundwater monitoring selected as the most viable remedial action alternative. A 2-year monitoring program implemented, after which it was decided that groundwater monitoring alone was insufficient to remediate the site.
Soil gas survey at OU-4	Phase II sampling (Law, 1989).	Law Environmental conducted a soil gas survey at 40 shallow soil-gas monitoring stations. The survey results indicated an elongated area of soil contamination stretching in a northwest/southeast direction and intersecting the fire training pit.
Reevaluate the proposed remedial actions	Draft RI/FS (Law, 1991)	Monitoring program results indicated possible contaminant migration in the shallow aquifer from the source area to Long Branch Creek. No surface water or sediment samples exhibited elevated site contaminants. Recommended the use of collection systems to halt contaminated groundwater before migrating to Long Branch Creek. Site officially named under the Resource Conservation and Recovery Act (RCRA) and officially named OU-4 under a separate Interim RCRA Facility Assessment (RFA) report.
Address remaining data gaps	Supplemental RI/FS work plan (SEC Donohue, Inc., 1992)	Performed topographic and geophysical surveys of the study area; recommended soil quality assessment in soil gas "hot spots"; installed additional monitoring wells (MW-116, -116A, -112A, -115A, -117, -118, and -119A); evaluated the hydrologic characteristics of the shallow aquifer; collected additional surface water and sediment sampling; performed a soil vapor extraction (SVE)/bioremediation treatability study; and performed data validation.
Perform soil SVE pilot test and bench scale bioremediation study	Draft Final Soil Vapor Extraction and Bioremediation Study Report (Rust E&I, 1993)	SVE pilot test and bioremediation study performed to aid in the selection of soil remediation alternatives for OU-4 as part of the RI/FS. SVE pilot test results indicated that the zone of influence for both vertical and horizontal SVE systems was limited. The bench-scale bioremediation study was performed on contaminated vadose soils at the site. The study indicated significant microbial activity on the basis of oxygen uptake and carbon dioxide evolution; however total petroleum hydrocarbon (TPH) and diesel-range organics (DRO) concentrations did not appear to decrease. Water and oxygen addition (e.g., through bioventing) were recommended for remediation of vadose soils on the basis of the bioremediation study.

# TABLE 1.1 (Concluded) HISTORY OF PAST REMEDIAL ACTIVITIES AT OU-4

# OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

		SHAW ML BOOTH CHANGE
Activity	Report/Reference	Results/Recommendations
Perform soil bioventing pilot test within fire training pit	Initiated 11/18/94; Streamlined test (AFCEE, 1996)	Bioventing test results indicate that TRPH and BTEX compounds were reduced between 1 to 3 orders-of-magnitude at the site. Successful hydrocarbon reductions through bioventing prompted the recommendation to continue bioventing at the discretion of the Base. The bioventing system continues to operate at the time of this report.
Perform chemical oxidation pilot test	Initiated 11/11/96; (IT, 1997)	Chemical oxidation pilot test performed to determine if groundwater contamination at OU-4 can be treated and reduced using hydrogen peroxide and ferrous iron (Fenton's reagent). Results indicated the generally uniform reductions of aromatic and chlorinated aliphatic hydrocarbons within an estimated effective treatment radius of 30 feet from the injection well. Off-gas of oxygen by <i>in situ</i> chemical reactions, dilution effects, and soil heterogeneity complicated quantifying the remedial effect of chemical oxidation. A full-scale chemical oxidation treatment system was recommended.
Groundwater pump and treat	None available.	At the time of this report, a groundwater pump-and-treat system is planned for the site as an interim measure (IM) under the direction of IT Corporation in an attempt to minimize plume migration and discharge to Long Branch Creek while the option of monitored natural attenuation is evaluated.



L: \45037\DRAWNGS\97DN0384, 01/15/98 ot 1:37

Beginning with the supplemental RI activities at the fire training area, the applicability of several remedial alternatives has been evaluated through pilot tests and bench-scale studies. The majority of these activities have focused on soils in the vicinity of the former (bermed) burn pit. Rust E&I (1993) conducted short-duration soil vapor extraction (SVE) pilot tests in the former bermed burn pit in 1993 to collect data for full-scale design, to monitor oxygen-enhanced respiration rates of indigenous microorganisms, and to compare the operating effectiveness of horizontal SVE wells versus vertical SVE wells. Success was limited due to vent well construction designs that contributed to air short-circuiting and soil water recovery (Rust E&I, 1993). A laboratory bench-scale study, also performed in 1993, concluded that a significant population of indigenous microorganisms is present in contaminated site soils, and rapid growth was exhibited by the microbes when supplied with oxygen (Rust E&I, 1993). In 1994, Parsons ES (1994 and 1995) performed a bioventing pilot test in a portion of the former bermed burn pit unaffected by the SVE system. After 1 year, soil concentrations of benzene, toluene, ethylbenzene, and xylenes (BTEX) decreased 1 to 3 orders of magnitude within approximately 30 feet of the air injection vent wells (Parsons ES, 1996). The bioventing pilot system has remained in operation in order to continue remediation of the heavier hydrocarbons in soils within the former bermed burn pit.

The most recent pilot test, conducted in November 1996, addressed chlorinated solvent contamination dissolved in groundwater in the vicinity of monitoring well MW-117, where the highest historical concentrations of dissolved chlorinated solvents have been detected at OU-4 [International Technology Corporation (IT), 1997]. The pilot test attempted to stimulate chemical oxidation through the one-time injection of hydrogen peroxide, sulfuric acid, and ferrous sulfate. The study concluded that approximately 80 percent of mass removal was attributed to stripping, with the remaining 20 percent attributed to chemical oxidation. This rate of chemical oxidation was approximately 35 percent of the anticipated chemical oxidation rate. Once again, these estimates were based on results collected one week after injection, and information regarding the long-term chemical oxidation effects is currently unavailable. The cost to remediate site groundwater using hydrogen peroxide injection was estimated at approximately 2.5 million dollars (IT, 1997).

### 1.5 SITE INVESTIGATION REPORTS

The results of previous site investigations are presented in the following reports:

- Installation Restoration Program, Phase I: Records Search, Shaw AFB, South Carolina (ES, 1983);
- Installation Restoration Program, Phase II: Confirmation 1 Quantification Stage 1, Shaw AFB, South Carolina (RTI, 1986);
- Remedial Action Plan, Shaw Air Force Base (Law, 1989);
- Draft Remedial Investigation and Feasibility Study Report for Site 1 Former Fire Training Area No. 1, Shaw AFB, Sumter, SC. Prepared for the US Army Corps of Engineers (Law, 1991);

- Draft Soil Vapor Extraction and Bioremediation Study Report (RUST E&I, 1993):
- Bioventing Test Work Plan for IRP Site FT-01 (Former Fire Training Area 1) and IRP Site SS-15 (POL Fuel Depot), Shaw AFB (Parsons ES, 1994);
- Draft Interim Bioventing Pilot Test Results Report, IRP Site FT-01 (Former Fire Training Area 1) and IRP Site SS-15 (POL Fuel Depot) Shaw AFB, SC (Parsons ES, 1995);
- Soil Vapor Extraction and Bioremediation Study Report, Operable Unit #4 (Rust E&I, 1993);
- Final RI Report, Operable Unit #4, Former Fire Training Area No. 1, IRP Site No. FT-1, Shaw AFB, SC (Rust E&I, 1995);
- Memorandum to Mr. Randy Adams, 20 CES/CEV, Kelly AFB, Re: Completion of Final Bioventing Test, Shaw AFB, Site FT-01 (Former Fire Training Area 1) and Site SS-15 (POL Fuel Depot) (Parsons ES, 1996);
- Final Feasibility Study/Corrective Measures Study Report and Proposed Plan, Operable Unit #4, Former Fire Training Area No. 1 (IRP Site No. FT-1, SWMU#59) (Rust E&I, 1996); and
- Draft Chemical Oxidation Pilot Test Report, Operable Unit 4, IRP Site Number FT-01, Shaw AFB, SC, Contract No. DACW 45-93-D-0044, TERC Region 2, Delivery Order 0004, AF Project No. VLSB-95-7002 (IT, 1997).

The site-specific data presented in Sections 3, 4, and 5 are based on a review of these documents and on data collected by Parsons ES under this program in April and May 1997.

### **SECTION 2**

### SITE CHARACTERIZATION ACTIVITIES

To meet the requirements of the RNA TS, additional data were required to evaluate near-surface geology and hydrogeology, contaminant distribution, and groundwater geochemistry. Site characterization activities involved borehole advancement and groundwater monitoring point installation with a cone penetrometer or Geoprobe®; collection of groundwater samples from existing monitoring wells and newly installed monitoring points; collection of surface water and sediment samples from adjacent creeks; and performance of slug tests on select groundwater monitoring wells. The scope of these activities is described in the work plan for this RNA TS (Parsons ES, 1997), and the work was performed in April and May 1997.

The physical and chemical data listed below were collected during the field work phase of the TS:

- Depth from measurement datum to the water table or potentiometric surface in monitoring wells and points;
- Stratigraphy of subsurface media;
- Groundwater geochemical data, including pH, temperature, electrical conductivity, total alkalinity, oxidation-reduction potential (ORP), dissolved oxygen (DO), carbon dioxide, chloride, nitrate+nitrite as nitrogen (N), ammonia, ferrous iron, manganese, sulfate, total organic carbon (TOC), dissolved hydrogen, and methane/ethane/ethene;
- Concentrations of CAHs, aromatic volatile organic compounds (VOCs) (including chlorinated benzenes), and metals in groundwater;
- Concentrations of CAHs and aromatic VOCs in surface water.

The following sections describe the procedures that were followed when collecting site-specific data. Additional details regarding investigative activities are presented in the work plan (Parsons ES, 1997).

### 2.1 CONE PENETROMETER TESTING

### 2.1.1 Monitoring Point Placement and Investigation of Site Stratigraphy with CPT

A cone penetrometer was used to install monitoring points at five locations at OU-4, identified as monitoring points MPA through MPE, on April 3 and 4, 1997.

Monitoring MPE was abandoned due to incorrect placement and reinstalled in May 1997 (using the same designation) with a Geoprobe® (Section 2.2) approximately 125 feet northeast of its former location. The CPT activities were conducted by the US Army Corps of Engineers (USACE). The cone penetrometer consists of an instrumented probe that is forced into the ground using a hydraulic load frame mounted on a heavy truck, with the weight of the truck providing the necessary reaction mass. The penetrometer equipment is housed in a stainless steel, dual-compartment body mounted on a 43,000-pound, triple-axle Kenworth® truck chassis powered by a turbo-charged diesel engine. The weight of the truck and equipment is used as ballast to achieve the overall push capability of 39,000 pounds.

The penetrometer probe has a 1.8-inch outside-diameter (OD), 60-degree conical point with sacrificial tip, and an 8.0-inch-long by 1.8-inch-OD friction sleeve. Inside the probe, two load cells independently measure the vertical resistance against the conical tip and the side friction along the sleeve. Each load cell is a cylinder of uniform cross-section that is instrumented with four strain gauges in a full-bridge circuit. As the probe is pushed into the subsurface, resistive forces are sensed by the load cells, and the data are transmitted from the probe assembly via a cable running through the push tubes. The analog data are digitized, recorded, and plotted by computer in the penetrometry truck to provide soil stratigraphy data.

After collecting soil stratigraphy data or placing a monitoring point (Section 2.3) with the cone penetrometer, the push rods were cleaned during extraction with the CPT steam-cleaning system (rod cleaner) as they were withdrawn from the ground. A vacuum system located beneath the cone penetrometer truck was used to recover cleaning water. Use of this system resulted in nearly 100-percent recovery of steam-cleaning rinseate from the rod cleaner. Precautions were taken to minimize any impact to the surrounding area that might result from decontamination operations. Potable water used in CPT equipment cleaning and decontamination was obtained from the Base water supply.

### 2.1.2 Investigation of Residual and Mobile Hydrocarbons

The known propensity of aromatic hydrocarbons to fluoresce under ultraviolet wavelengths has allowed the use of LIF technology, in conjunction with CPT technology, to evaluate soil characteristics and hydrocarbon contamination simultaneously. The LIF system has a 0.25-inch sapphire window in the side of the cone that allows a nitrogen laser to scan the soil for fluorescent compounds as the LIF penetrometer rod pushes through soil. The wavelength used in the USACE CPT/LIF system gives the strongest fluorescence signal (attributable to the presence of contamination) for naphthalene and heavier, long-chained hydrocarbons. Thus, while the LIF is not entirely appropriate for detecting the fluorescence of BTEX, it is useful for defining soil contamination because the heavy long-chained hydrocarbons are more likely to sorb to the soil matrix than the more soluble BTEX compounds. Fiber optic cables connected to the laser spectrometer and a 6-pair electrical conductor connected to the CPT data acquisition system are routed through the interior of the push tubes to the CPT probe.

Graphical results of each LIF push are provided in Appendix A. The graphs showed cone resistance, sleeve friction, soil classification, fluorescence intensity, and maximum fluoresced wavelength.

### 2.2 GEOPROBE® FIELD ACTIVITIES

Geoprobe® field work occurred on May 15 through May 17, 1997. The truck-mounted Geoprobe® system was used to re-install monitoring point MPE, originally installed at an incorrect location using the CPT. The Geoprobe® system is a hydraulically-powered percussion/probing machine used to advance sampling tools through unconsolidated soils. The system relies on static weight from the vehicle combined with percussion as the energy for advancement of the sampling tool. Geoprobe® push rods were manually driven into the ground where the Geoprobe® truck did not have access to install six temporary monitoring points (TMP-1 through TMP-6). TMP-6 did not produce water at a screened depth of 10 feet bgs after two installation attempts and was abandoned.

### 2.3 GROUNDWATER MONITORING POINT COMPLETION

The term "monitoring point" is used in this report to distinguish these groundwater monitoring stations from conventionally constructed monitoring wells. Completion details for new monitoring points and previously installed monitoring wells are summarized in Table 2.1 and detailed in Appendix A. Figure 2.1 shows the locations of newly completed monitoring points. There are currently 21 groundwater monitoring wells, 3 temporary monitoring wells, and 3 piezometers associated with OU-4. As part of the current investigation, 5 CPT monitoring points and 6 Geoprobe® temporary monitoring points were installed at 11 locations.

### 2.3.1 Monitoring Point Construction

### 2.3.1.1 Pre-Installation Activities

All subsurface utility lines and other man-made subsurface features were located, and proposed monitoring point locations were cleared and approved by the Base prior to any drilling activities. Monitoring point locations were adjusted as necessary to avoid damage to subsurface utilities.

### 2.3.1.2 Equipment Decontamination Procedures

Prior to arriving at the site and between each monitoring point location, all Geoprobe® rods, tips, sleeves, pushrods, samplers, tools, and other downhole equipment were decontaminated using an Alconox® detergent and potable water solution and scrub brush, followed by a potable water rinse.

### 2.3.1.3 Monitoring Point Materials Decontamination

Monitoring point completion materials were inspected by the field scientist and determined to be clean and acceptable prior to use. All monitoring point completion materials were factory sealed in plastic wrap. Pre-packaged casing, sand, bentonite,

TABLE 2.1
SUMMARY OF WELL INSTALLATION DETAILS
OU-4 RNA TS
SHAW AFB, SOUTH CAROLINA

Well	Installation	Northing	Easting	Casing Diameter	TOC Elevation	Ground Elevation	Total Well Depth	Screen Length
Identification	Date	(feet)	(feet)	(inches)	(ft msl) <sup>a/</sup>	(ft msl)	(ft bgs) <sup>b/</sup>	(feet)
	3/8/93	782128.419	2166707.029	2.00	214.87	212.92	19.70	5.00
MW-105 MW-105D	3/8/93 7/13/88	782128.419 782134.801	2166707.029	2.00	214.87	212.92	176.50	10.00
MW-105D	3/10/93	782452.344	2166601.959	2.00	219.21	217.63	24.30	5.00
MW-106D	7/21/88	782432.344	2166595.258	2.00	218.99	217.03	179.50	10.00
MW-100D MW-107	3/10/93	782281.684	2166296.107	2.00	214.79	213.21	19.90	5.00
MW-107 MW-111	3/11/93	782234.008	2165610.262	2.00	227.08	225.35	27.60	5.00
MW-111 MW-112	3/9/93	782254.008	2166456.628	2.00	218.01	215.78	25.00	15.00
MW-112A	2/4/93	782153.806	2166465.881	2.00	217.92	215.78	89.70	10.33
MW-112A MW-112D	7/24/88	782159.916	2166459.506	2.00	217.73	215.78	155.20	10.33
MW-112D	3/10/93	782139.910	2166473.268	2.00	208.95	206.79	25.00	15.00
MW-113A	3/10/93	782490.479	2166472.454	2.00	209.22	207.07	35.00	15.00
MW-114	3/8/93	782369.261	2166797.623	2.00	207.10	204.83	25.00	15.00
MW-115	3/11/93	781886.414	2166877.511	2.00	204.19	201.92	25.00	15.00
MW-115A	2/18/93	781888.874	2166869.608	2.00	204.19	201.92	69.08	10.31
MW-116	2/1/93	781981.943	2166241.078	2.00	219.46	217.20	25.60	15.30
MW-116A	1/21/93	781982.286	2166251.067	2.00	219.48	217.14	94.74	10.33
MW-117	2/5/93	781914.192	2166673.356	2.00	216.42	213.89	21.41	15.24
MW-118	2/15/93	781610.669	2166979.110	2.00	203.51	201.12	10.36	10.31
MW-119A	5/17/94	782159.953	2167543.911	2.00	205.89	203.59	75.30	10.33
TW-101	11/9/86	781707.969	2166693.132	4.00	220.47	218.08	60.00	30.00
TW-120	4/97	782021.000	2167099.000	2.00	194.64	NA <sup>c</sup>	5.30	NA
TW-121	4/97	781842.000	2167099.000	2.00	194.04	NA NA	3.30 NA	NA NA
PZ-101	11/8/86	781665.342	2166718.029	2.00	NA	218.53	30.00	10.00
PZ-101 PZ-102	11/11/86	781686.963	2166705.655	2.00	NA NA	218.53	28.00	10.00
PZ-102	11/11/86	781680.303	2166741.982	2.00	NA NA	218.63	35.00	10.00
MPA	4/3/97	NA	NA	0.50	NA NA	NA	18.00	9.84
MPB	4/3/97	NA NA	NA NA	0.50	NA NA	NA NA	18.00	9.84
MPC	4/3/97	NA	NA	0.50	NA	NA	14.00	9.84
MPD	4/3/97	NA	NA	0.50	NA	NA	23.00	9.84
				SS <sup>d</sup>				
MPE	5/16/97	NA	NA		NA	NA	20.00	9.84
TMP-1	5/15/97	NA	NA	SS	NA	NA	8.00	0.50
TMP-2	5/15/97	NA	NA	SS	NA	NA	6.50	0.50
TMP-3	5/15/97	NA	NA	SS	NA	NA	6.50	0.50
TMP-4	5/15/97	NA	NA	SS	NA	NA	8.00	0.50
TMP-5	5/16/97	NA	NA	SS	NA	NA	6.00	0.50
TMP-6	5/16/97	NA	NA	SS	NA	NA	8.00	0.50

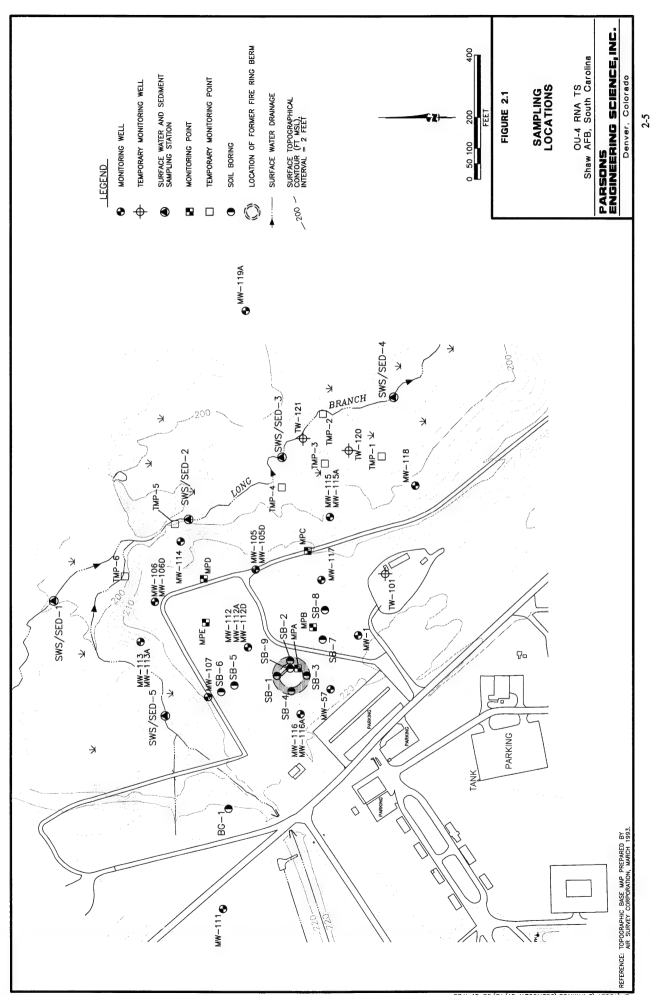
a/ ft msl = feet above mean sea level.

Note: Survey data for monitoring points MPA through MPE were incomplete, therefore indicated as NA.

 $<sup>^{</sup>b/}$  ft bgs = feet below ground surface.

o' NA = Not available.

 $<sup>^{</sup>d'}$  SS = Constructed of 0.25-inch-ID stainless steel screen.



and concrete mix were used in point construction, and the bags were inspected for possible external contamination before use. Materials that could not be cleaned to the satisfaction of the field scientist were not used.

### 2.3.1.4 Monitoring Point Casing and Screen

Two monitoring point designs were used to construct shallow monitoring points. The four monitoring points (MPA through MPD) installed using the CPT were constructed of Schedule 40 polyvinyl chloride (PVC) riser pipe connected to factory-slotted PVC screens having an inside diameter (ID) of 0.5 inch. All PVC casing and screen sections on the shallow monitoring points were flush threaded; glued joints were not used. The riser pipe at each PVC monitoring point was fitted with a PVC top cap, and an aluminum drive point was inserted into the bottom of each PVC screen. The one monitoring point (MPE) installed using the hydraulically-powered Geoprobe® apparatus and the six temporary monitoring points (TMP-1 through TMP-6) driven manually into the ground were constructed with 0.25-inch-ID stainless steel (SS) mesh implants acting as monitoring point screens and 0.25-inch-ID, Teflon®-lined, high-density polyethylene (HDPE) tubing acting as risers connecting the SS mesh to the surface. The SS mesh implants were attached to an aluminum drive points.

Monitoring point screens constructed of PVC were approximately 10 feet long and factory-slotted with 0.010-inch openings. The monitoring points constructed with Teflon®-lined HDPE tubing had screens constructed of 0.5-foot-long SS mesh with pore openings of 0.0057 inch. The bottoms of the SS mesh screens were threaded to dedicated aluminum drive points/implant anchors that remained in place after the drive rods were removed, leaving the riser tubing extended to the surface. The field scientist verified and recorded the borehole depth and the lengths of all casing sections and tubing. All lengths and depths were measured to the nearest 0.1 foot. All monitoring points were screened near the water table.

### 2.3.2 Monitoring Point Installation

Using the CPT truck, USACE staff installed four 0.5-inch-ID PVC monitoring points MPA, MPB, MPC, and MPD within OU-4 (Figure 2.1). The evacuated hole remaining after extracting the pushrods was cased with PVC riser pipe (introduced through an internal tube running the length of the push rod assembly) as the rods were withdrawn and steam cleaned. These monitoring points were screened at intervals between 4 and 23 feet below ground surface (bgs). Prior to the installation of any CPT monitoring points, soil stratigraphy data were collected with the CPT device from the ground surface to 18 to 50 feet bgs. CPT stratigraphy data are included in Appendix A.

Using the hydraulically powered Geoprobe® apparatus, USEPA staff installed a single 0.2-inch-ID monitoring point at MPE within OU-4 by driving lengths of 1.0-inch-OD probe rod to the desired depth of 20 feet bgs. The downhole end of the drive rod was prefitted with a sacrificial aluminum drive point to decrease soil resistance and to prevent soils from entering the drive rod. After reaching the desired depth, a stainless steel mesh implant was connected to HDPE tubing and threaded through the

center of the Geoprobe® drive rod. The drive rods were then removed with the hydraulically-powered apparatus, leaving the sacrificial tip, screen, and riser tubing in the formation at the desired depth. Formation soils quickly collapsed into the borehole after the removal of the drive rod to create a natural sand pack. Because soils collapsed to the ground surface, no annular sealant (e.g., bentonite) could be placed around the tubing.

Parsons ES staff installed six temporary monitoring points TMP-1 through TMP-6 (Figure 2.1) by manually driving lengths of 1.0-inch-OD probe rod to the desired depth with a 30-pound rod driver. The drive rod was prefitted with a sacrificial aluminum drive point, and HDPE tubing was threaded into this point in the same manner described for MPE. The drive rods were then manually removed, leaving the sacrificial tip, SS screen, and HDPE riser tubing in the formation at the desired depth. The depths to screens ranged from 6 to 8 feet bgs. Formation soils quickly collapsed into the boreholes after the removal of the drive rod to create a natural sand pack. As a result of soils quickly collapsing to the ground surface, no annular sealants (e.g. bentonite) could be placed as a seal around the tubing.

#### 2.3.3 Monitoring Point Development

Prior to sampling, newly installed monitoring points were developed. Typically, well development removes sediment from inside the well casing and flushes fines, cuttings, and drilling fluids from the sand pack and the portion of the formation adjacent to the well screen. Use of the CPT and Geoprobe® to place monitoring points eliminates cuttings and drilling fluids. Therefore, development of these monitoring points was primarily intended to minimize the amount of fine sediment that might accumulate in the casing. Clayey soils in the vicinity of TMP-6 prevented successful monitoring point development after two attempts, and this point was abandoned.

Development was accomplished for monitoring points MPA through MPD using a peristaltic pump with dedicated silicone and HDPE tubing. The pump tubing was regularly lowered to the bottom of the PVC points so that fines were agitated and removed from the point in the development water. For monitoring point MPE and all temporary monitoring points, development was performed by attaching the peristaltic pump tubing directly to the HDPE riser tubing. Typically, development was continued until a minimum of 10 casing volumes of water were removed, and the pH, temperature, and conductivity of the groundwater had stabilized. Development waters were containerized and disposed of at the Base industrial wastewater treatment plant (IWTP). Development records are included in Appendix A.

#### 2.4 GROUNDWATER SAMPLING

This section describes the procedures used for collecting groundwater samples. In order to maintain a high degree of quality control (QC) during this sampling event, the procedures described in the TS work plan (Parsons ES, 1997) and summarized in the following sections were followed. A summary of the chemical analyses performed for groundwater, as well as surface water, sediment, and mobile LNAPL, is presented in Table 2.2.

#### **TABLE 2.2** ANALYTICAL PROTOCOL USED FOR GROUNDWATER, SURFACE WATER, SEDIMENT, AND MOBILE LNAPL SAMPLES

#### **OU-4 RNA TS** SHAW AFB, SOUTH CAROLINA

MATRIX Analyte	METHOD	FIELD (F) OR FIXED-BASE LABORATORY
GROUNDWATER		
Ferrous Iron (Fe <sup>2+</sup> )	Colorimetric, Hach Method 8146	F
ORP	Direct-reading meter	F
Oxygen	Direct-reading meter	F
Dissolved Hydrogen	Reduction Gas Analyzer	F
Hydrogen Sulfide	Colorimetric, Hach Method 8131	
pH	Direct-reading meter	F
Conductivity	Direct-reading meter	F
Temperature	Direct-reading meter	F
Carbon Dioxide	CEHMetrics Method 4500	F
Alkalinity (Carbonate [CO <sub>3</sub> <sup>2</sup> ] and Bicarbonate [HCO <sub>3</sub> ])	F = Titrimetric, Hach Method 8221	F
Nitrate + Nitrite	USEPA Method 353.1	NRMRL <sup>a/</sup>
Chloride	N-601 <sup>b/</sup>	NRMRL
Sulfate	N-601	NRMRL
Methane, Ethane, Ethene	RSKSOP-175/RSKSOP-147°	NRMRL
Ammonia-Diss. Gas in Water	E350.1	NRMRL
Total Organic Carbon Aromatic Hydrocarbons	RSKSOP-102	NRMRL
+ Fuel Carbon	RSKSOP-122	NRMRL
Chlorinated Volatile Organics	RSKSOP-148	NRMRL
URFACE WATER		
Aromatic Hydrocarbons		NRMRL
+ Fuel Carbon	RSKSOP-122	
EDIMENT		
VOCs	USEPA Method 525.2, Revision 3	NRMRL
TOC	RSKSOP-102	NRMRL
MOBILE LNAPL <sup>d</sup>		
VOCs	USEPA Method 525.2, Revision 3	NRMRL

 <sup>&</sup>lt;sup>a/</sup> USEPA National Risk Management Research Laboratory (NRMRL) in Ada, Oklahoma.
 <sup>b/</sup> Waters capillary electrophoresis Method N-601.

<sup>&</sup>lt;sup>c'</sup> RSKSOP = Robert S. Kerr Laboratory (now NRMRL) Standard Operating Procedure.

d LNAPL = Light nonaquous-phase liquid.

#### 2.4.1 Groundwater Sampling Locations

Groundwater samples were collected from 17 previously installed monitoring wells, and 10 of the 11 newly installed monitoring points. Monitoring point TMP-6 could not be developed, and was abandoned. After completion of installation and development activities, the monitoring points were purged and sampled using a peristaltic pump with dedicated HDPE and silicone tubing for monitoring points MPA through MPD, and by directly attaching the peristaltic pump to the HDPE riser for monitoring point MPE and temporary monitoring points TMP-1 through TMP-5.

#### 2.4.2 Preparation and Equipment Cleaning

All equipment used for sampling was assembled and properly cleaned and calibrated (if required) prior to use in the field. Calibrations were performed in accordance with the manufacturer's specifications. An electrical water level meter was used to measure the static water level in the monitoring well/point prior to initiation of purging. Prior to each use, the water level probe was cleaned with a potable water and phosphate-free, laboratory-grade detergent solution, followed by a distilled-water rinse. In addition, a clean pair of new, disposable latex or nitrile gloves was worn each time a different well or monitoring point was sampled. Dedicated HDPE and silicone tubing were used at each sampling location, eliminating the need for decontaminating these items between wells.

#### 2.4.3 Groundwater Sampling Procedures

#### 2.4.3.1 Preparation of Location

Prior to sampling, the area around the well or monitoring point was cleared of foreign materials, such as brush, rocks, and debris. These procedures prevented sampling equipment from inadvertently contacting debris around the monitoring well/point. The integrity of the monitoring well/point also was inspected, and any irregularities in the visible portions of the well/point, protective cover, or concrete pad were noted.

#### 2.4.3.2 Water Level and Total Depth Measurements

Prior to removing any water from the well or monitoring points, an electrical water level probe or oil/water interface probe was used to measure the depth to groundwater below the well datum to the nearest 0.01 foot. Water levels in the newly installed points constructed of Teflon®-lined HDPE tubing were not obtained because the water level probe diameter was larger than the tubing diameter. If the monitoring well/point depth was not known, the water level probe was then lowered to the bottom of the well/point for measurement of total depth (recorded to the nearest 0.1 foot). Based on these measurements, or using total depths recorded in the RI report (Rust E&I, 1995), the volume of water to be purged from the well/point was calculated. Water level data are reviewed in Section 3.

#### 2.4.3.3 Monitoring Well/Point Purging

Where possible, a minimum of three times the calculated saturated casing volume was removed from each monitoring well/point prior to sampling. Purging continued until the pH, DO concentration, ORP, conductivity, and temperature stabilized between successive readings. Physical and chemical parameters were measured at the well head using field meters and a flow-through cell consisting of an Erlenmeyer flask. peristaltic pump with dedicated silicone and HDPE tubing was used for well evacuation at all monitoring points and wells TW-120, TW-121, MW-105, MW-111, MW-112, MW-117, and MW-118. For the monitoring points with 0.25-inch HDPE risers, purging was accomplished by attaching the silicone peristaltic pump tubing directly to the top of the monitoring point tubing. For monitoring points with PVC risers and for monitoring wells purged with the peristaltic pump, the HDPE tubing was lowered to within 2 feet of the bottom of the casing. A two-stage submersible pump was used to purge wells MW-107, MW-113, MW-114, MW-115, MW-116, and MW-116A; a onestage submersible pump was used to purge MW-106; and a Bennett pump was used to purge wells MW-112A and TW-101. In each case, the positive pressure pump was lowered to within 2 feet of the bottom of the casing. The pumps were decontaminated between wells with Alconox® and a distilled-water rinse. TMP-6 did not purge when screened at a depth of 10 feet bgs, and therefore was not sampled. A replacement for TMP-6 was installed at 8 feet bgs, and also did not purge. All purge water was containerized and disposed of at the Base IWTP. Purging and sampling field forms are presented in Appendix A.

#### 2.4.3.4 Sample Collection

A peristaltic pump with dedicated silicone and HDPE tubing was used to extract groundwater samples from the same wells where it was used for purging, along with TW-101, MW-106, MW-112A, and MW-116. In wells containing mobile LNAPL, the pump tubing was lowered through the LNAPL layer to the groundwater under positive air flow to prevent contamination of the inside of the tubing. A two-stage pump was used for sampling wells MW-107, MW-113, MW-114, MW-115, and MW-116A. In almost all cases, the sampling was performed immediately following well purging. For example, at wells where a peristaltic pump was used, the pump was not turned off between purging and sampling activities. In a few instances, the monitoring well/point was purged dry, and the samples were collected after sufficient recharge had occurred. All samples were collected within an hour of purging.

The samples were transferred directly into the appropriate sample containers. The water was carefully poured down the inner walls of each sample bottle to minimize aeration of the sample. Sample bottles for aromatic and chlorinated VOCs and dissolved gases (methane and ethene) were filled so that there was no headspace or air bubbles within the container.

#### 2.4.4 Onsite Chemical Parameter Measurement

DO measurements were taken using a flow-through cell at the outlet of the sampling pump. DO concentrations were recorded after the readings stabilized, and in all cases

represent the lowest DO concentration observed. Because the electrical conductivity, pH, ORP, and temperature of the groundwater can change significantly within a short time following sample acquisition, these parameters also were measured in the field, in the same flow-through cell used for DO measurements. Measured values were recorded on groundwater sampling records (Appendix A). Table 2.2 lists the field methods used to analyze the individual parameters, and analytical results are presented in Section 4.

#### 2.4.5 Sample Handling

The fixed-base analytical laboratory, the National Risk Management Research Laboratory (NRMRL), provided pre-preserved sample containers where appropriate. The sample containers were filled as described in Section 2.4.3.4, and the container lids were tightly closed. The samples were labeled as described in the work plan.

After the samples were sealed and labeled, they were transported to the onsite USEPA mobile laboratory where several analyses were performed. Analytical results are presented in Section 4. USEPA personnel packaged the samples for fixed-base laboratory analysis to prevent breakage and leakage or vaporization from the containers. Sample shipment to NRMRL and the associated chain-of-custody documentation was the responsibility of the USEPA/NRMRL field personnel.

#### 2.5 SURFACE WATER SAMPLING

Surface water samples were collected from five locations along Long Branch Creek and its tributaries in the northeastern portion of OU-4 (Figure 2.1). These samples were collected in order to assess the degree to which VOC-contaminated groundwater was discharging to the surface drainages. The surface water samples were analyzed for both total fuel carbon and CAHs (Table 2.2).

Surface water samples were collected from the west side of Long Branch Creek by placing the sample bottle in the channel with the opening facing up and allowing the water to slowly fill the bottle. Sample handling proceeded as described for groundwater samples in Section 2.4.5. Analytical results are discussed in Section 4.

#### 2.6 SEDIMENT SAMPLING

Stream sediment samples were collected from the same five locations within Long Branch Creek as the surface water samples (Figure 2.1, Table 2.2) following surface water collection. These samples were collected to assess the degree to which VOCs have adsorbed to the bottom of the stream channel. The sediment samples were analyzed for VOCs and TOC (Table 2.2), and results are presented in Section 4.

Sediment samples were collected using a hand auger to excavate 8 inches into the stream bed. The samples were then transferred from the auger to sample containers. Sample handling proceeded as described for groundwater samples in Section 2.4.5.

#### 2.7 LNAPL SAMPLING

Mobile LNAPL samples were collected to assess the chemical composition of on-site source contamination. Free product was encountered at temporary monitoring points MPB and MPC, and sampled using a peristaltic pump. Product samples were collected after groundwater sampling, as the LNAPL level fell during groundwater extraction. If recharge from the aquifer maintained the level of the LNAPL, the product was sampled by slightly raising the extraction tubing.

Mobile LNAPL samples were pumped directly into sample containers. Sample handling proceeded as described for groundwater samples in Section 2.4.5. The analytical method for mobile LNAPL samples is listed in Table 2.2, and results are discussed in Section 4.

#### 2.8 AQUIFER TESTING

During the RI, Rust E&I (1995) performed a pump test at TW-101, and transmissivities were calculated using both the Theis type curve and semilog methods of analysis at TW-101, PZ-101, PZ-102, PZ-103, MW-115, and MW-117. For comparison, slug tests were performed at wells MW-106 and MW-112 to provide estimates for hydraulic conductivity of the shallow aquifer. Slug tests are single-well hydraulic tests used to estimate the hydraulic conductivity of an aquifer in the immediate vicinity of the tested well. Slug testing can be performed using either a rising head or a falling head test. Both rising head and falling head tests were used at this site. Detailed slug testing procedures are presented in the Technical Protocol for Implementing Intrinsic Remediation with Long-Term Monitoring for Natural Attenuation of Fuel Contamination Dissolved in Groundwater (Wiedemeier et al., 1995), hereafter referred to as the Technical Protocol document.

Slug tests were performed using a Hermit® data logger and a slug of PVC pipe capped at each end. Data obtained during slug testing were analyzed using the computer program AQTESOLV® (Geraghty & Miller, Inc., 1994) and the methods of Bouwer and Rice (1976) and Bouwer (1989) for unconfined conditions. The results of slug testing are presented in Section 3.3.2.2 and Appendix A.

#### **SECTION 3**

#### PHYSICAL CHARACTERISTICS OF THE STUDY AREA

This section describes the physical characteristics of OU-4 as determined from data collected by Parsons ES in April and May 1997, in conjunction with pertinent data documented in previous site reports (see Section 1.5).

#### 3.1 TOPOGRAPHY, SURFACE HYDROLOGY, AND CLIMATE

Shaw AFB lies along the border of the Upper and Middle Coastal Plain subprovinces of the Atlantic Coastal Plain Physiographic Province. This transitional boundary is marked by the Orangeburg scarp and expressed at the surface by a change in elevation (Rust E&I, 1995).

Surface water and drainage features in the vicinity of the Base are shown on Figure 3.1. The small perennial stream, Long Branch Creek, and an unnamed southwest drainage ditch both border the Base on the east and south, respectively. The east-flowing drainage ditch joins the south-flowing Long Branch Creek southeast of the Base, approximately 1 mile downstream from Booth's Pond. Several other ponds, including Sawmill Pond, Frierson Pond, and Loring Mill Pond, also are located southeast of the Base (Figure 3.1).

Topography surrounding OU-4 can be characterized as relatively flat in the vicinity of the former bermed fire pit to steeply sloping near Long Branch Creek. Long Branch Creek and an ephemeral tributary that enters Long Branch Creek from the west occupy the Long Branch Creek flood plain and mark the eastern and northern site boundaries, respectively. Elevations range from approximately 198 feet above mean sea level (msl) at Long Branch Creek to approximately 216 feet msl at the bermed former burn pit. Surface runoff drains to the north and east into the bordering streams (Rust E&I, 1995).

In the vicinity of OU-4, Long Branch Creek is 8 to 10 feet wide, 2 to 4 feet deep, and flows at an estimated rate of 1,000 gallons per minute. Long Branch Creek is impounded to form Booth's Pond approximately 2,000 feet downstream from the site. When flowing, the ephemeral stream along the northern boundary of OU-4 can have a width of approximately 4 feet and a depth of 9 inches.

The mean annual temperature at Shaw AFB is 64 degrees Fahrenheit (°F), with an average of 46 °F in January and 81 °F in July. Precipitation averages approximately 46 inches per year distributed relatively evenly throughout the year, with the highest monthly maximum in July and the lowest in October. Average relative humidity is

relatively constant throughout the year, ranging from 72 to 86 percent (Rust E&I, 1995).

#### 3.2 REGIONAL GEOLOGY AND HYDROGEOLOGY

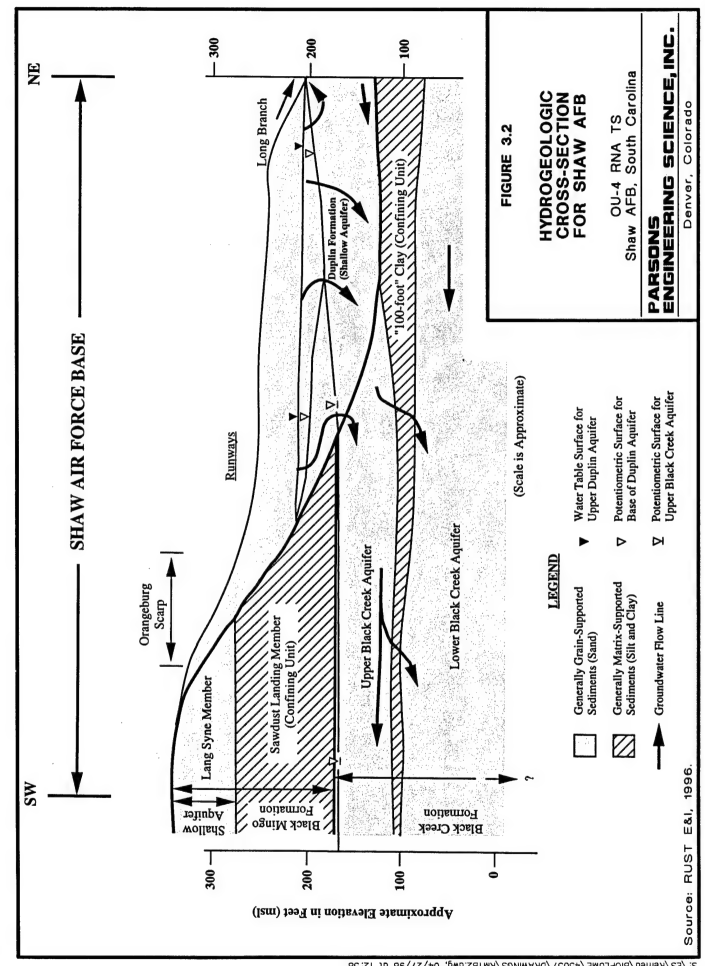
Shaw AFB is underlain by Cretaceous to Quaternary sediments and sedimentary rocks that in turn overlie a basement of Paleozoic crystalline and Triassic sedimentary rocks. These sediments and sedimentary rocks are present as a sedimentary wedge that thickens and dips to the south and southeast. The sedimentary rock sequence at Shaw AFB is approximately 600 to 700 feet thick. Cretaceous formations in this area make up the greatest thickness of sediments and are overlain by 100 to 200 feet of Tertiary and Quaternary sediments. The sedimentary geologic units at the Base, described in ascending order (decreasing age) are the Middendorf Formation, the Black Creek Formation, the Black Mingo Formation, and the Duplin Formation (Figure 3.2).

The Cretaceous Black Creek Formation is the deepest formation encountered to date during environmental investigations at OU-4 (Figure 3.2). The lithology of the Black Creek Formation is predominantly fine- to medium-grained, interbedded, white and gray sands with thin sandstone beds and gray, black, and dark-blue clays. The sediments contain fossils, including shells, sharks teeth, and microfauna. Intervals may contain pyrite, pyrite-coated wood fragments, and marcasite, and may be phosphatic, glauconitic, and micaceous, with interbedded, finely laminated organic clays.

The Paleocene-age Black Mingo Formation (Figure 3.2) outcrops west of the runway at Shaw AFB, and is composed of glauconitic quartz sands, thin layers of gray to light-green silty clay, pyritic dark-gray organic clays, and Fuller's earth (opaline claystone). The Black Mingo Formation locally includes two members: the Sawdust Landing Member and the Lang Syne Member. The Sawdust Landing Member consists of sands, silts, and clays representing upper delta plain deposits, similar to the modern Mississippi River Delta. In the Sawdust Landing Member, significant clay and silt content represents flood-stage deposits, while fine to coarse sands represent channel deposits. As the Paleocene sea transgressed the land, it laid down the cleaner Lang Syne sands and clays. The Lang Syne represents the beach barrier island and back-barrier depositional environments at the seaward margin of the Sawdust Landing delta.

At Shaw AFB, the Duplin Formation unconformably overlies the Black Mingo and Black Creek Formations, and outcrops throughout the eastern two-thirds of the Base. The Orangeburg Scarp is expressed at the ground surface where the Pliocene-age Duplin Formation contacts the eroded surface of the underlying deposits. Locally, this scarp is expressed as the hill traversing the western edge of the Base along Highway 441 (Figure 1.2). The Duplin Formation consists largely of sands, silts, and clays that were derived from the underlying formations and reworked during a sea-level decrease.

Shaw AFB is underlain by three aquifer systems: the Middendorf Aquifer System, the Black Creek Aquifer System, and the Shallow Aquifer System. Aquifer system boundaries do not necessarily correspond to boundaries between geologic formations. In addition, lithologies may change significantly from one area to another within these aquifer systems. The Black Creek and Shallow Aquifer Systems are described below



because they are used for the Base water supply and their quality has been adversely impacted by Base activities. A schematic representation of the local aquifers is presented in Figure 3.2, which demonstrates the conceptual relationships among the various geologic formations, aquifer systems, and the Orangeburg Scarp.

The Black Creek Aquifer System underlies most of Sumter County and is used for water supply over most of the central Coastal Plain, including for most of the Base supply. In the vicinity of Shaw AFB, the Black Creek Aquifer is present between approximately 175 feet above msl and 25 feet below msl, and is separated into a lower and upper aquifer by a clay aquitard referred to as the 100-foot clay (Figure 3.2).

In the western portion of the Base, the Shallow Aquifer System is present in sediments associated with the Black Mingo Formation; however, in the vicinity of OU-4, the aquifer is present in the Duplin Formation. The Shallow Aquifer extends from the water table (commonly at 210 to 240 feet above msl) to approximately 150 to 175 feet above msl.

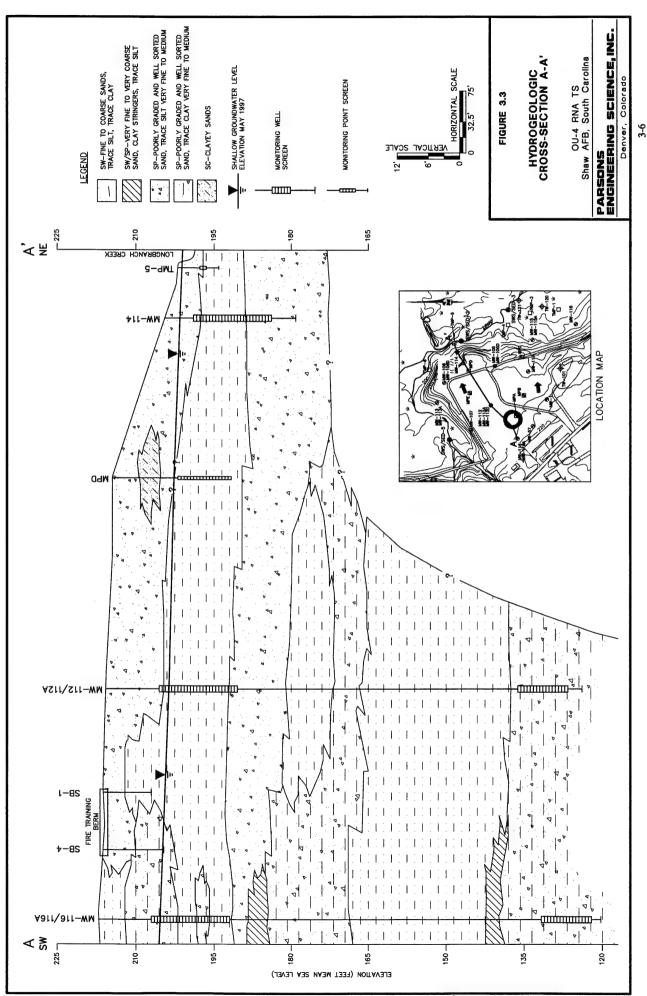
#### 3.3 SITE GEOLOGY AND HYDROGEOLOGY

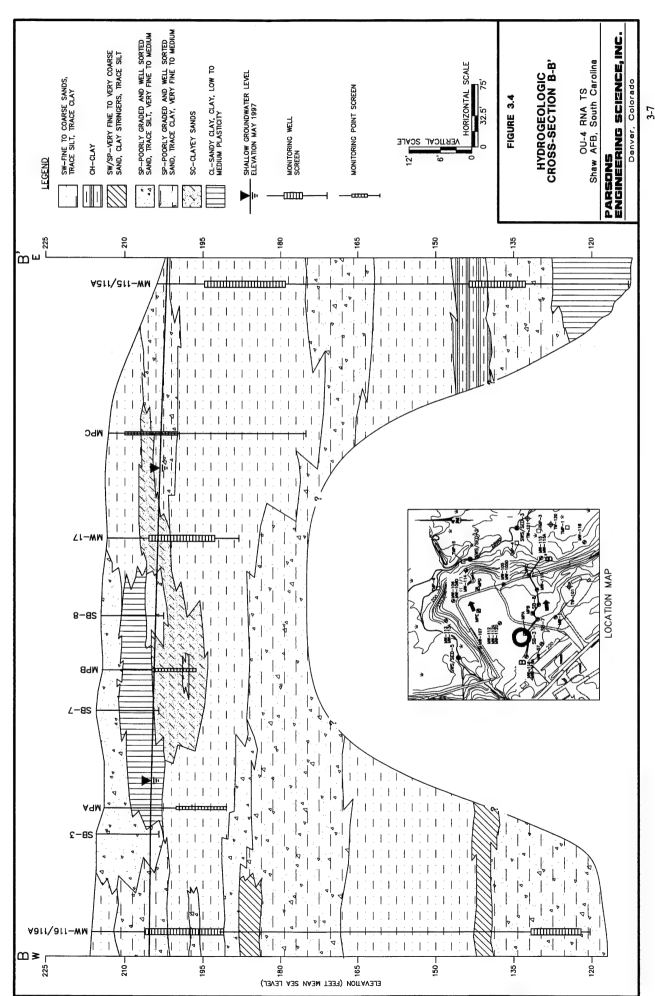
Two cross-sections modified from the final RI (Rust E&I, 1995) with geologic data collected during the TS field activities, are shown on Figures 3.3 and 3.4. Both cross-sections are roughly parallel to groundwater flow and contaminant migration paths to the northeast and east-southeast.

#### 3.3.1 Lithology and Stratigraphic Relationships

OU-4 lies approximately 1 mile east of the surface expression of the Orangeburg Scarp; therefore, site geology consists of Duplin Formation terrace deposits directly underlain by the Black Creek Formation. The Duplin Formation extends from the surface to a depth of about 80 to 90 feet bgs. Overall, the Duplin Formation at OU-4 consists of unconsolidated, well to poorly graded sands, with approximately 15 percent interstitial clays and silts. Soils in the upper part of this unit are mostly coarse- to fine-grained sands; however, a thin, discontinuous clayey unit was observed in the shallow subsurface in a number of site borings. The lower part of the Duplin Formation consists mostly of medium to fine sands, with clay content increasing with depth. Clay is present both interstitially and in seams. The transition between the lower and upper portions of this unit is gradational.

At OU-4, upper portions of the Black Creek Formation were eroded prior to the deposition of the Duplin Formation terrace deposits, so that the first encountered unit is a clay referred to as the "100-foot clay." This clay unit aquitard was encountered at 90 to 110 feet bgs in three OU-4 boreholes (MW-106D, MW-112D, MW-115D), and is about 20 to 35 feet thick. The unit consists of very stiff, light-gray and very pale brown, plastic clay with seams of poorly graded sands, sandy clays, and silts. The continuity of the silt seams is uncertain. Previous OU-4 investigations indicate that site contamination has not penetrated this clay unit; therefore, the reader is referred to the final RI report (Rust E&I, 1995) for descriptions of the deeper site geology.





The water-bearing zone of primary interest at the site is the Shallow Aquifer. At OU-4, this aquifer coincides with the Duplin Formation terrace deposits. The upper portion of the aquifer is characterized by mostly medium to very coarse sands with gravelly layers, all containing interstitial fine sands and clays. With increasing depth, sediments gradually become finer. At a depth of approximately 90 feet bgs, the "100-foot clay" of the Black Creek Formation acts as an aquitard between the shallow aquifer and the underlying Lower Black Creek Aquifer. This is indicated by large head differences between the shallow aquifer and Lower Black Creek Aquifer.

A localized unit consisting of low-permeability, clayey sand is present in the area of the former fire pit at depths ranging from 3 to 9 feet bgs. This low-permeability unit results in both a local groundwater perching effect after heavy rain and a local confining effect on the underlying groundwater. With the exception of this clayey unit, the Shallow Aquifer is unconfined. As the clayey unit was not observed in all borings, its continuity and areal extent have not been fully delineated. Its occurrence at varying depths suggests that the layer may not be continuous.

#### 3.3.2 Groundwater Hydraulics

In the Shallow Aquifer, depth to groundwater ranges from approximately 20 feet bgs beneath the upland terrace to a few inches bgs near Long Branch Creek. May 1997 groundwater elevations are shown in Table 3.1.

#### 3.3.2.1 Flow Direction and Gradient

The shallow aquifer is markedly influenced by surface topography. This is reflected by relatively shallow gradients within the upland terrace portion of the site, and steeper gradients parallel to and underlying the embankments along the flood plain of Long Branch Creek and the unnamed ephemeral stream. As shown on Figure 3.5, groundwater generally flows toward the east/northeast in the shallow aquifer at OU-4. Horizontal groundwater gradients computed from May 1997 data average approximately 0.0048 foot/foot (ft/ft) in the upland terrace portion of the site near the former fire training pit. The groundwater gradient increases to approximately 0.011 ft/ft on the terrace slope and near the creek. The average groundwater gradient from the former fire training pit to the Long Branch Creek is 0.007 ft/ft.

Long Branch Creek receives groundwater from OU-4 through natural gravitational migration of groundwater down the upland terrace and along the embankments of the creek. The creek becomes a hydraulic sink for groundwater (and groundwater contaminants) and acts as a downgradient hydraulic boundary for groundwater migration. This observation is supported by Strack (1989) who provides an example in which a stream penetrates one-tenth of the aquifer thickness and captures approximately 94 percent of the groundwater flow from its ugradient side. The creek depth penetrates as much as five feet into the aquifer in the vicinity of OU-4, where contaminants are primarily limited to the upper 30 feet of shallow aquifer (Section 4). Therefore, the majority of groundwater/dissolved contamination reaching the creek probably will discharge into it with a small fraction hydraulically capable of migration past the creek.

# TABLE 3.1 SUMMARY OF GROUNDWATER ELEVATION DATA MAY 1997 OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

	Elevation of		
	Reference Point	Depth to	Groundwater
Well	for Measurements	Groundwater	Elevation
Identification	(feet msl) <sup>a/</sup>	(feet btc) <sup>b/</sup>	(feet msl)
MW-105	214.87	13.54	201.33
MW-106	219.21	17.79	201.42
MW-107	214.79	11.43	203.36
MW-111	227.08	21.05	206.03
MW-112	218.01	15.06	202.95
MW-112A	217.92	19.71	198.21
MW-113	208.95	7.19	201.76
MW-113A	209.22	7.46	201.76
MW-114	207.10	7.28	199.82
MW-115	204.19	4.52	199.67
MW-115A	204.16	7.18	196.98
MW-116	219.46	15.28	204.18
MW-116A	219.48	21.06	198.42
MW-117	216.42	14.73	201.69
MW-118	203.51	4.06	199.45
TW-101	220.47	18.73	201.74

<sup>&</sup>lt;sup>a/</sup> Feet msl = feet above mean sea level.

b/ Feet btc = feet below top of casing.



The horizontal gradients measured in May 1997 are consistent with the horizontal gradients measured in March 1993 that ranged from 0.006 ft/ft to 0.033 ft/ft. However, observed heads from March 1993 generally were 2 feet higher. On the basis of groundwater elevations from four wells screened nearer the bottom of the shallow aquifer, the lower zone of the Shallow Aquifer has a horizontal groundwater gradient of 0.003 ft/ft to the southeast.

In addition to differences in flow direction, static water level measurements indicate that there are head differences between the upper and lower zones of the Shallow Aquifer. Downward vertical hydraulic gradients between the upper and lower zones were 0.077 ft/ft at well cluster MW-116, 0.071 ft/ft at well cluster MW-112, and 0.050 ft/ft at well cluster MW-115.

#### 3.3.2.2 Hydraulic Conductivity

As a part of supplemental RI field activities, Rust E&I (1995) performed a 26-minute pump test at well TW-101, which is screened from 30 to 60 feet bgs and is located approximately 400 feet southeast of the fire training pit (Figure 2.1), transverse to the direction of contaminant migration. The average hydraulic conductivity measured at this location was 17.6 feet per day (ft/day) (Rust E&I, 1995). In May 1997, Parsons ES conducted falling- and rising-head slug tests at wells MW-106 and MW-112 and analyzed the test data using the method of Bouwer and Rice (1976) (see Section 2.8). These slug tests yielded hydraulic conductivities of 81 ft/day and 68 ft/day (average of 75 ft/day) measured at wells MW-106 (screened from 14.9 to 19.9 feet bgs and MW-112 (screened from 10 to 25 feet bgs), respectively. These wells are located along the northeastern contaminant migration flow path. The range of hydraulic conductivities measured from the pump test and slug tests (18 to 81 ft/day) are indicative of silty to clean sands.

#### 3.3.2.3 Effective Porosity

Because of the difficulty involved in accurately determining effective porosity, published literature values for soil types comprising the shallow saturated zone were referenced (Spitz and Moreno, 1996). Estimates of effective porosity for fine to medium sand range from 0.08 to 0.30. An average effective porosity of 0.25 was assumed for the Shallow Aquifer at OU-4 considering the predominance of medium-grained sands in the upper saturated zone of the aquifer.

#### 3.3.2.4 Advective Groundwater Velocity

The advective velocity of groundwater in the direction parallel to groundwater flow is given by:

$$\overline{v} = \frac{K}{n_o} \frac{dH}{dL}$$

Where:  $\bar{\nu} = \text{Average advective groundwater velocity } [L/T]$  K = Average slug-test hydraulic conductivity [L/T] (75 ft/day) dH/dL = Average gradient [L/L] (0.007 ft/ft)  $n_e = \text{Effective porosity } (0.25).$  Using this relationship in conjunction with site-specific data, the average advective groundwater velocity at site OU-4 in May 1997 was 2.1 ft/day, or approximately 770 feet per year (ft/yr).

#### 3.4 GROUNDWATER USE

Potential receptor exposure to groundwater contamination could result through use of water obtained from on-Base supply wells screened within the surficial and Black Creek aquifers. The closest supply well (Base Well 8) is screened within the Shallow Aquifer approximately 700 feet upgradient from the site near the Munitions Flight Unit parking area. There are no other supply wells screened within the Shallow Aquifer in the vicinity of OU-4. The closest supply well screened in the Black Creek Formation is Base Well 10, located approximately 2,000 feet southwest of (upgradient from) OU-4 (Rust E&I, 1996). Base Well 10 reportedly has been abandoned (Rust E&I, 1995). The absence of producing groundwater wells in the vicinity of OU-4 reduces the possibility of a groundwater exposure pathway to potable sources. Off-Base migration of dissolved contaminants to the east/northeast is not expected, because shallow groundwater migrating through OU-4 discharges to Long Branch Creek.

#### **SECTION 4**

### CONTAMINANT DISTRIBUTION AND EVIDENCE FOR BIODEGRADATION

Soil and groundwater have been contaminated as a result of fire training activities at OU-4. Work performed as part of the RFA (Law, 1989), the RI/FS (Law, 1991; SEC Donohue, Inc., 1992), and supplemental RI work (Rust E&I, 1994) for OU-4 focused on defining the nature and extent of contamination. Data collected during those efforts were supplemented with data collected during natural attenuation sampling in May 1997 and are presented in the following subsections. In particular, this section focuses on data useful for evaluating and modeling the natural attenuation of fuel and chlorinated hydrocarbons dissolved in groundwater.

#### 4.1 CONTAMINANT SOURCES AND SOIL QUALITY

#### 4.1.1 Mobile LANPL Contamination

During fire training exercises between 1941 and 1969, numerous contaminants were burned during fire training exercises, including jet fuel, waste oil, hydraulic fluid, spent solvents, contaminated fuel, and napalm. It is believed that some of the metal drums used to store and transport the combustibles may have been buried at the site, subsequently contributing to soil and groundwater contamination (Rust E&I, 1995). Mobile LNAPL (i.e., free-phase product) was not detected during the previous RFA, RI/FS, or RI investigations; however, mobile LNAPL was observed at monitoring points MPB, MPC, and MPD in May 1997. Groundwater from MPD exhibited only a sheen, while thicker product layers were encountered at MPB and MPC. LNAPL samples were collected from MPB and MPC for laboratory analysis. LNAPL at monitoring point MPB was brown in color, highly aromatic, and had a viscosity similar to gasoline when shaken. The LNAPL observed at monitoring point MPC was dark, less aromatic than the LNAPL measured at MPB, and appeared to be more viscous than gasoline when shaken. Several inches of LNAPL likely was present at monitoring points MPB and MPC, though the thicknesses of the LNAPL layers could not be measured because the diameters of the monitoring points (0.5-inch ID) were smaller than the diameter of the oil/water interface probe and available bailers.

The analytical results for product samples collected at MPB and MPC are shown in Table 4.1. The LNAPL samples appear to be extremely weathered, especially with regard to benzene. The benzene, toluene, ethylbenzene, and xylenes (BTEX) information in Table 4.1 is insufficient to identify the original fuel type, but the weathered LNAPL samples likely originated from JP-4 or AVGAS releases. The cosolvenation of trichloroethene (TCE) in the MPC LNAPL suggests that this LNAPL source may have resulted from simultaneous burning of fuels and solvents during

#### TABLE 4.1 MOBILE LNAPL RESULTS OU-4 RNA TS SHAW AFB. SOUTH CAROLINA

Compound	Concentration in Fresh JP-4 a/ (mg/L) c/	Concentration in Product from MPB b/ (mg/L)	Concentration in Product from MPC (mg/L)
Benzene	3,750	ND d/	26.4
Toluene	9,975	1,422	3,170
Ethylbenzene	2,775	847	1,140
Total Xylenes	17,400	3,662	4,823
TCE	NA <sup>e/</sup>	ND	3,350
Total BTEX	33,900	5,931	9,159.4

<sup>&</sup>lt;sup>a</sup> Data from Martel (1987).

previous fire training operations or leakage from drums previously stored or currently buried at the site.

#### 4.1.2 Soil Contamination

Numerous soil samples were collected during previous site investigations to characterize the extent of soil contamination. Additional soil samples were not collected during this TS; however, LIF data obtained as a part of CPT activities provide additional information on the extent of soil contamination. The areal extent of total VOC contamination in soils as defined during the RI and modified using LIF data is presented on Figure 4.1. A summary of VOC and diesel-range organics analysis results for RI soil samples collected in February/March 1993 is provided in Table 4.2. Detected VOC concentrations consisted primarily of fuel constituents; however, traces

of 1,1,1-trichloroethane (TCA), 1,2-dichloroethene (DCE), and tetrachloroethene (PCE) were each measured at four locations.

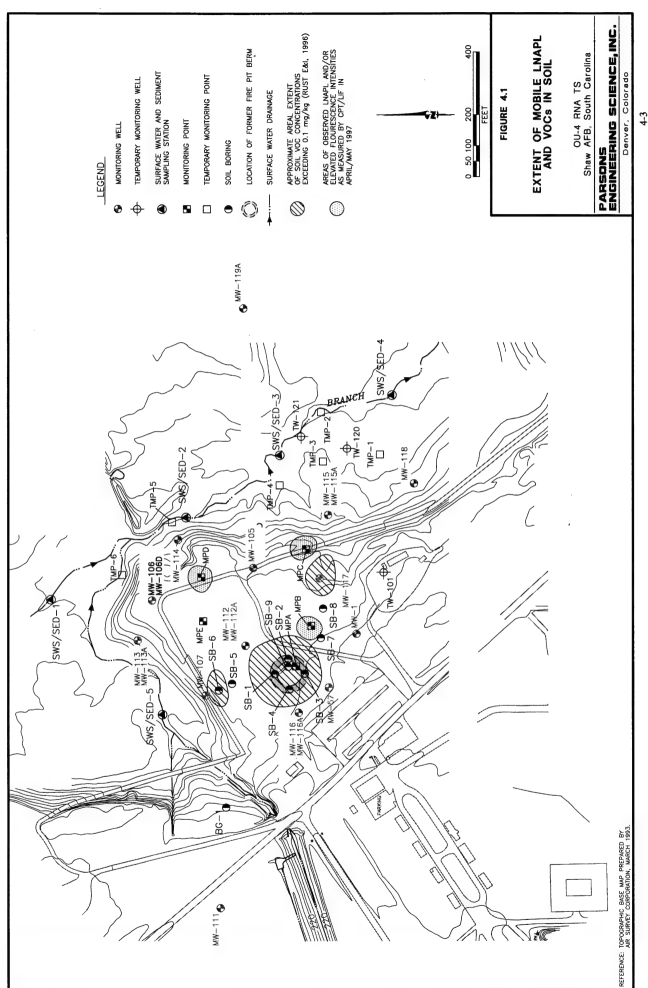
The highest total VOC concentrations measured during the RI were identified in soil samples collected from a depth of 6 to 8 feet bgs. VOC concentrations exceeding 100 micrograms per kilogram (µg/kg) [0.1 milligrams per kilogram (mg/kg)], centered on three areas that roughly coincided with the three "hot spots" identified from a soil gas survey performed at the site (Law, 1989). The highest total VOC concentrations in soil were detected in the vicinity of the former bermed burn pit at concentrations of up to

by USEPA Method 525.2, Revision 3.

c' mg/L = Milligrams per liter.

d ND = Not detected.

e/ NA = Not applicable.



**TABLE 4.2** SUMMARY OF DETECTED VOLATILE AND DIESEL-RANGE ORGANIC COMPOUNDS IN OU-4 SOIL SAMPLES COLLECTED IN FEBRUARY/MARCH 1993 **OU-4 RNA TS** SHAW AFB, SOUTH CAROLINA

			Total			Total	Diesel-Range
Sample	1,1,1-TCA	PCE	1,2-DCE	Ethylbenzene	Toluene	Xylenes	Organics
ID	$(\mu g/kg)^{a/}$	$(\mu g/kg)$	(µg/kg)	(µg/kg)	$(\mu g/kg)$	(µg/kg)	(mg/kg) <sup>b/</sup>
MW-112A-2	< 5.6°	3.3 J <sup>d/</sup>	< 5.6	< 5.6	< 5.6	< 5.6	<3.5
MW-112A-8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	<3.8
MW-112A-14	< 5.7	< 5.7	< 5.7	< 5.7	< 5.7	< 5.7	14
MW-115A-3	< 6.8	< 6.8	11	5 J	6.5 J	14	47
MW-116-2	< 5.2	< 5.2	< 5.2	< 5.2	< 5.2	< 5.2	5.4
MW-116-7	< 5.2	< 5.2	< 5.2	< 5.2	< 5.2	< 5.2	4.1
MW-116-12	< 5.7	< 5.7	< 5.7	< 5.7	< 5.7	< 5.7	4.6
MW-116A-2	< 5.2	2.6 J	< 5.2	< 5.2	< 5.2	< 5.2	6.2
MW-116A-8	< 5.5	< 5.5	< 5.5	< 5.5	< 5.5	< 5.5	<3.7
MW-116A-14	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	<3.8
MW-117-2	< 5.4	< 5.4	< 5.4	< 5.4	< 5.4	< 5.4	13
MW-117-7	< 30	< 30	< 30	100	< 30	640	190
MW-117-12	< 5.9	< 5.9	< 5.9	4.3 J	< 5.9	9.4	5.5
MW-118-2	< 6.3	< 6.3	< 6.3	< 6.3	< 6.3	< 6.3	13
MW-119A-1	< 5.4	< 5.4	< 5.4	< 5.4	< 5.4	< 5.4	3.9
MW-119A-4	< 5.6	< 5.6	< 5.6	< 5.6	< 5.6	< 5.6	2.3J
SB-1-2	< 28	< 28	< 28	150	28	440	7300
SB-1-8	< 570	< 570	< 570	3000	< 570	10000	6300
SB-1-12	< 5500	< 5500	< 5500	17000	23000	75000	5200
SB-2-2	< 5.3	< 5.3	< 5.3	< 5.3	< 5.3	< 5.3	200
SB-2-8	< 570	490 J	< 570	140 J	< 570	630	1200
SB-3-2	< 5.5	< 5.5	< 5.5	< 5.5	< 5.5	< 5.5	20
SB-3-8	< 550	< 550	< 550	150J	< 550	750	3500
SB-3-12	< 56	< 56	< 56	190	< 56	1500	430
SB-4-2	< 530	< 530	< 530	490 J	170 J	3700	9500
SB-4-8	< 14000	< 14000	< 14000	< 14000	< 14000	35000	5700
SB-4-12	< 6000	< 6000	< 6000	10000	13000	91000	2000
SB-5-2	< 27	< 27	< 27	< 27	< 27	< 27	550
SB-5-8	< 5.9	< 5.9	< 5.9	< 5.9	< 5.9	< 5.9	<3.8
SB-5-12	< 5.7	< 5.7	< 5.7	< 5.7	< 5.7	< 5.7	8
SB-6-2	< 5.5	< 5.5	< 5.5	< 5.5	< 5.5	< 5.5	290
SB-6-8	140 J	< 570	< 570	1900	300 J	12000	1100
SB-6-12	< 5600	< 5600	< 5600	4700 J	1300 J	23000	110
SB-7-2	< 5.3	< 5.3	< 5.3	< 5.3	< 5.3	< 5.3	81
SB-7-8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	5.4
SB-7-12	< 5600	< 5600	< 5600	< 5600	< 5600	< 5600	650
SB-8-2	< 5.3	< 5.3	< 5.3	< 5.3	< 5.3	< 5.3	5.4
SB-8-8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	< 5.8	<3.4
SB-9-2	< 530	< 530	< 530	1300	< 530	3100	3900
SB-9-6	< 560	< 560	< 560	640	< 560	980	3600
SB-9-10	< 570	< 570	< 570	790	< 570	710	3400

<sup>&</sup>quot;ug/kg = Micrograms per kilogram.

Source: Rust E&I, 1995.

b'mg/kg = Milligrams per kilogram.

a'<= Not detected down to 10% of the quantitation limit listed.

b'J = Estimated value less than the quantitation limit.

115,000  $\mu g/kg$  (115 mg/kg as combined total of ethylbenzenes, toluene, and xylenes). This area also contained the only VOC concentrations detected above 100  $\mu g/kg$  in soil samples collected from 0 to 3 feet bgs (at SB-4 and SB-9). Nevertheless, the detection of VOCs in two areas beyond the still-recognizable burn pit suggested that secondary contaminant sources are present at the site. These two areas center around MW-117, east of the pit, and SB-6 located north of the pit (Figure 4.1).

Soil and mobile LNAPL contamination detected during this TS expanded the known extent of soil contamination identified during the RI, especially in the vicinity of MW-117. Fluorescent aromatic compounds indicative of hydrocarbon contamination were observed at locations MPA, MPB, and MPC (also coinciding with mobile LNAPL observations at MPB and MPC) during monitoring point installation with the CPT/LIF. As shown on Figure 4.1, mobile LNAPL (Section 4.1.1) and LIF data obtained at monitoring point MPB indicate a possible connection between soil contamination measured in the fire training pit and around MW-117. LNAPL may have migrated away from the training pit toward MW-117; however, mobile LNAPL likely originated from two separate sources, one near the fire training pit (MPB) and the other near a suspected former drum storage or fire training location (MPC). Similarly, mobile LNAPL and LIF data obtained at monitoring point MPC suggests that soil contamination extends farther north/northeast of MW-117 than delineated during the RI.

Although relatively thin zones of fluorescent compounds were identified atop the water table during CPT/LIF activities, it is possible that an LNAPL smear zone exists elsewhere in the vicinity of MW-117 that contributes to mobile LNAPL volumes as groundwater elevations fluctuate. The detection of high fluorescence intensities from 8 to 12 feet bgs at MPC suggests that mobile LNAPL is residing on the water table and acting as the primary contaminant source at MPC [the groundwater depth at MPC is approximately 10 feet bgs (Table 3.1)]. High fluorescence intensities from 6 to 15 feet bgs at monitoring point MPB suggest the possible presence of residual LNAPL contamination (in addition to the mobile LNAPL) at this location and may indicate a zone of smeared contamination above and below the water table. Fluorescence detections from 6 to 10 feet bgs at MPA in the fire training pit are consistent with previous detections of VOCs in soil in this area (Table 4.2), and indicate that significant fuel contamination might still be present. CPT/LIF logs are reproduced in Appendix A.

Soil contamination profiles measured during the RI most likely have been altered through site remediation activities that have occurred over the past 3 years. For example, after 1 year of bioventing in the former bermed burn pit, BTEX concentrations decreased more than 90 percent within approximately 30 feet of the vent wells (Parsons ES, 1996b). The effects of bioventing beyond 30 feet from the vent wells and after an additional 1.5 years of bioventing are not known. Also, hydrogen peroxide injection during November 1996 in the vicinity of MW-117 may have induced changes in soil VOC concentrations in this area. During the pilot test, water levels were temporarily raised to the ground surface, which may have contributed to an increase in the thickness of the residual LNAPL smear zone, resulted in *in situ* soil washing, and/or caused the accelerated migration of mobile LNAPL (LNAPL was observed at MPC, downgradient from MW-117). Soil gas emissions also increased during the test as a result of groundwater sparging; it is unclear if soil gas emissions

affected soil VOC concentrations. Remediation that could have affected soil VOC concentrations in the vicinity of the soil contamination near SB-6 has not been attempted.

The areal extent of DRO concentrations roughly coincides with the extent of VOC contamination (Table 4.2); however, DROs were more widespread than total VOCs in 0- to 3-foot bgs samples. This is a probable result of the site grading reported by Rust E&I (1995). The disturbance from grading likely would volatilize any VOCs in shallow soils while dispersing the DRO over a somewhat larger area. Given that the berm for the former burn pit is still evident, it is likely that this area was not graded; therefore, any VOCs residing in the shallow soils in this area would have remained undisturbed.

#### 4.2 OVERVIEW OF HYDROCARBON BIODEGRADATION

Mechanisms for natural attenuation of chlorinated solvents and BTEX include biodegradation, dispersion, dilution from recharge, sorption, and volatilization. Of these processes, biodegradation is the only mechanism working to transform contaminants into ultimately innocuous byproducts. Intrinsic bioremediation occurs when indigenous microorganisms work to bring about a reduction in the total mass of contamination in the subsurface without the addition of nutrients. In order to provide a foundation for interpreting site data, the following subsections review the major bioremediation processes that act upon CAHs and BTEX. Because the terminology describing biodegradation of both types of compounds is similar, a generalized review of the processes is presented first.

#### 4.2.1 Review of Biodegradation Processes

Microorganisms obtain energy for cell production and maintenance by facilitating thermodynamically advantageous reduction oxidation (redox) reactions involving the transfer of electrons from electron donors to available electron acceptors. This results in the oxidation of the electron donor and the reduction of the electron acceptor. Electron donors may be natural organic carbon, fuel hydrocarbon compounds, and less-chlorinated solvents [e.g., vinyl chloride (VC), DCE, dichloroethane (DCA), or chlorinated benzenes ranging from chlorobenzene (CB) to tetrachlorobenzene (TeCB)]. Fuel hydrocarbons or solvents are completely degraded or detoxified if they are utilized as the primary electron donor (i.e., as a primary substrate or carbon source) for microbial metabolism (Bouwer, 1992). Electron acceptors are elements or compounds that occur in relatively oxidized states, and include oxygen, nitrate, ferric iron, sulfate, manganese, carbon dioxide, and highly chlorinated solvents [e.g., PCE, TCE, tetrachloroethane (PCA), TCA, and polychlorinated benzenes].

The driving force of biodegradation is electron transfer, which is quantified by the Gibbs free energy of the reaction ( $\Delta G^{\circ}_{r}$ ) (Stumm and Morgan, 1981; Bouwer, 1994; Godsey, 1994). The value of  $\Delta G^{\circ}_{r}$  represents the quantity of free energy consumed ( $\Delta G^{\circ}_{r} > 0$ ) or yielded ( $\Delta G^{\circ}_{r} < 0$ ) to the system during the reaction. Although thermodynamically favorable, most of the reactions involved in biodegradation of fuel hydrocarbons or chlorinated hydrocarbons cannot proceed abiotically because of the lack of activation energy. Microorganisms are capable of providing the necessary activation energy; however, they will facilitate only those redox reactions that have a

net yield of energy (i.e.  $\Delta G^{\circ}_{r} < 0$ ). Most reactions involving biodegradation of contaminants do yield energy to the microbes; however, in many cases specific geochemical conditions are necessary for this reaction to be favorable and to allow the appropriate microbial population to develop and grow.

Microorganisms preferentially utilize electron acceptors while metabolizing hydrocarbons (Bouwer, 1992). DO is utilized first as the prime electron acceptor. It is under these conditions (i.e., aerobic conditions) that fuel hydrocarbons and the less chlorinated solvents are most commonly used as electron donors. After the DO is consumed, anaerobic microorganisms use native electron acceptors in the following order of preference: nitrate, manganese, ferric iron hydroxide, sulfate, and finally carbon dioxide. Under anaerobic conditions, BTEX compounds and other fuel hydrocarbons are still used as electron donors. Chlorinated solvents that are susceptible to reductive dehalogenation generally are used as electron acceptors when aquifer conditions are such that sulfate or carbon dioxide is the preferred electron acceptor.

In addition to being controlled by the energy yield of the reaction, the expected sequence of redox processes is also a function of the ORP of the groundwater. This potential is a measure of the relative tendency of a solution or chemical reaction to accept or transfer electrons. As each subsequent electron acceptor is utilized, the groundwater becomes more reducing, and the ORP of the water decreases. The main force driving this change in ORP is microbially mediated redox reactions. ORP can be used as an indicator of which redox reactions may be operating at a site. Environmental conditions and microbial competition ultimately determine which processes will dominate.

Depending on the types and concentrations of electron acceptors present (e.g., nitrate, sulfate, carbon dioxide), pH conditions, and ORP, anaerobic biodegradation can occur by denitrification, manganese reduction, ferric iron reduction, sulfate reduction, or methanogenesis. Other, less common anaerobic degradation mechanisms such as manganese or nitrate reduction may dominate if the physical and chemical conditions in the subsurface favor use of these electron acceptors. Vroblesky and Chapelle (1994) show that the dominant terminal electron accepting process can vary both temporally and spatially in an aquifer with fuel hydrocarbon contamination.

#### 4.2.2 Biodegradation of BTEX

Numerous laboratory and field studies have shown that hydrocarbon-degrading bacteria can participate in the degradation of many of the chemical components of fuel hydrocarbons, including the BTEX compounds (Jamison et al., 1975; Atlas, 1981, 1984, 1988; Gibson and Subramanian, 1984; Reinhard et al., 1984; Young, 1984; Bartha, 1986; Wilson et al., 1986, 1987, and 1990; Barker et al., 1987; Baedecker et al., 1988; Lee, 1988; Chiang et al., 1989; Grbic-Galic, 1989 and 1990; Cozzarelli et al., 1990; Leahy and Colewell, 1990; Altenschmidt and Fuchs, 1991; Alvarez and Vogel, 1991; Baedecker and Cozzarelli, 1991; Ball et al., 1991; Bauman, 1991; Borden, 1991; Brown et al., 1991; Edwards & Grbic-Galic., 1992; Evans et al., 1991a and 1991b; Haag et al., 1991; Hutchins and Wilson, 1991; Hutchins et al., 1991a and 1991b; Beller et al., 1992; Bouwer, 1992; Edwards and Grbic-Galic, 1994; Thierrin et al., 1992; Malone et al., 1993; Davis et al., 1994). Fuel hydrocarbons biodegrade naturally when an indigenous population of hydrocarbon-degrading microorganisms is

present in the aquifer and sufficient concentrations of electron acceptors and nutrients are available to these organisms.

During biodegradation of fuel hydrocarbons, the fuel compounds are used as electron donors. They are the primary substrate ("food") for the microbes, while the electron acceptors provide the oxidant used to reduce (metabolize) the substrate and produce energy. In most subsurface environments, both aerobic and anaerobic degradation of fuel hydrocarbons can occur, often simultaneously in different parts of the plume. The nearly ubiquitous nature of these processes is well documented (e.g., Rice et al., 1995; Wiedemeier et al., 1995; Kuehne and Buscheck, 1996, and Mace et al., 1997). Aerobic destruction of BTEX compounds dissolved in groundwater results in the consumption (reduction) of DO and the formation of carbon dioxide. Anaerobic destruction of the BTEX compounds is associated with the accumulation of fatty acids, production of methane, solubilization of iron, and reduction of nitrate and sulfate (Cozzarelli et al., 1990; Wilson et al., 1990).

#### 4.2.3 Biodegradation of Chlorinated Solvents

Chlorinated solvents can be transformed, directly or indirectly, by biological processes (Bouwer et al., 1981; Miller and Guengerich, 1982; Reineke and Knackmuss, 1984; Wilson and Wilson, 1985; de Bont et al., 1986; Nelson et al., 1986; Spain and Nishino, 1987; Bouwer and Wright, 1988; Little et al., 1988; Mayer et al., 1988; Arciero et al., 1989; Cline and Delfino, 1989; Freedman and Gossett, 1989; Folsom et al., 1990; Harker and Kim, 1990; Alvarez-Cohen and McCarty, 1991a, 1991b; DeStefano et al., 1991; Henry, 1991; Sander et al., 1991; McCarty et al., 1992; Hartmans and de Bont, 1992; Chapelle, 1993; McCarty and Semprini, 1994; Vogel, 1994; Suflita and Townsend, 1995; Bradley and Chapelle, 1996; Klier et al., 1996; Spain, 1996). Biodegradation of CAHs and chlorinated benzenes results from the same general processes that bring about biodegradation of fuel hydrocarbons. However, a more complex series of processes often is involved, and CAHs and chlorinated benzenes may act as either substrates (electron donors) or electron acceptors, depending upon the prevailing geochemical conditions.

Whereas BTEX are biodegraded in essentially one step by acting as an electron donor/carbon source, CAHs and chlorinated benzenes may undergo several types of biodegradation involving several steps. Chlorinated solvents may undergo biodegradation through three different pathways: use as an electron acceptor, use as an electron donor, or cometabolism, which is degradation resulting from exposure to a catalytic enzyme fortuitously produced during an unrelated process. At a given site, one or all of these processes may be operating, although at many sites the use of CAHs as electron acceptors appears to be most likely.

In a pristine aquifer, native organic carbon is utilized as an electron donor and DO is utilized first as the prime electron acceptor. Where anthropogenic carbon (e.g., fuel hydrocarbons, less-chlorinated CAHs, or chlorinated benzenes with four or fewer chlorines) is present, it also may be utilized as an electron donor. Most chlorinated solvents that can act as electron donors have thus far only been demonstrated to do so under aerobic conditions, with the notable exception of VC (Bradley and Chapelle, 1996). After the DO is consumed, anaerobic microorganisms typically use native electron acceptors (as available) in the following order of preference: nitrate, ferric

iron oxyhydroxide, sulfate, and finally carbon dioxide. Evaluation of the distribution of these electron acceptors can provide evidence of where and how biodegradation of chlorinated solvents is occurring. In addition, because solvents may be used as electron acceptors or electron donors (in competition with other acceptors or donors), isopleth maps showing the distribution of these compounds also will provide evidence of the types of biodegradation processes acting at a site. A more complete description of the main types of biodegradation reactions affecting chlorinated solvents dissolved in groundwater is presented in the following subsections.

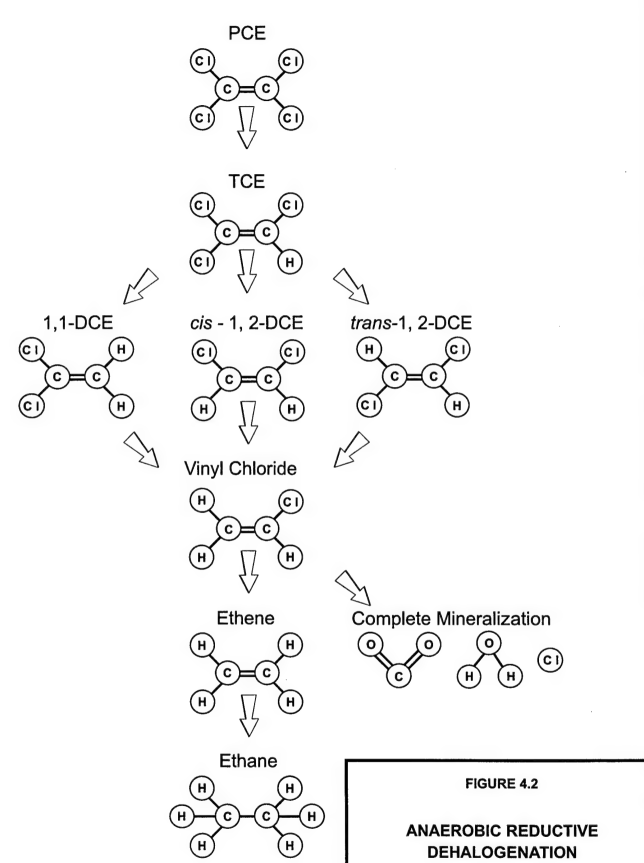
#### 4.2.3.1 Electron Acceptor Reactions (Reductive Dehalogenation)

Under anaerobic conditions, biodegradation of chlorinated solvents usually proceeds through a process called reductive dehalogenation. During this process, the halogenated hydrocarbon is used as an electron acceptor, not as a source of carbon, and a halogen atom is removed and replaced with a hydrogen atom. As an example, Figure 4.2 illustrates the transformation of chlorinated ethenes via reductive dehalogenation. In general, reductive dehalogenation of chlorinated ethenes occurs by sequential dehalogenation from PCE to TCE to DCE to VC to ethene. An analogous pattern for chlorinated ethanes might be PCA to TCA to DCA to chloroethen, and for chlorinated benzenes the pattern might be TeCB to trichlorobenzene (TCB) to dichlorobenzene (DCB) to CB. Depending upon environmental conditions, these sequences may be interrupted, with other processes (e.g., aerobic or abiotic degradation) then acting upon the products. Reductive dehalogenation of chlorinated solvent compounds is associated with the accumulation of daughter products and an increase in chloride.

Reductive dehalogenation affects chlorinated compounds differently. Of the ethenes, PCE is the most susceptible to reductive dehalogenation because it is the most oxidized. Conversely, VC is the least susceptible to reductive dehalogenation because it is the least oxidized of these compounds. Likewise, hexachlorobenzene is more susceptible to reductive dehalogenation than less-chlorinated benzenes. In general, the rate of reductive dehalogenation of chlorinated solvents has been observed to decrease as the degree of chlorination decreases (Vogel and McCarty, 1985; Bouwer, 1994). Murray and Richardson (1993) have postulated that this rate decrease may explain the accumulation of VC in PCE and TCE plumes that are undergoing reductive dehalogenation.

In addition to being affected by the degree of chlorination of the compound, reductive dehalogenation also can be controlled by the redox conditions of the site groundwater system. In general, reductive dehalogenation has been demonstrated under anaerobic nitrate- and sulfate-reducing conditions, but the most rapid biodegradation rates, affecting the widest range of compounds, occur under methanogenic conditions (Bouwer, 1994). Dehalogenation of PCE and TCE to DCE can proceed under mildly reducing conditions such as nitrate reduction or iron(III) reduction (Vogel et al., 1987), while the transformation of DCE to VC, or the transformation from VC to ethene requires more strongly reducing conditions (Freedman and Gossett, 1989; DeStefano et al., 1991; De Bruin et al., 1992).

Reductive dehalogenation of some compounds also has been shown to preferentially produce specific daughter compounds. For example, during reductive dehalogenation of TCE or PCE, all three isomers of DCE can theoretically be produced; however,



**OU-4 RNA TS** Shaw AFB, South Carolina

PARSONS ENGINEERING SCIENCE, INC.

Denver, Colorado

Bouwer (1994) reports that under the influence of biodegradation, *cis*-1,2-DCE is a more common intermediate than *trans*-1,2-DCE, and that 1,1-DCE is the least prevalent intermediate of the three DCE isomers.

When chlorinated compounds are used as electron acceptors, there must be an appropriate source of carbon for microbial growth in order for reductive dehalogenation to occur (Bouwer, 1994). Potential carbon sources/electron donors can include low-molecular-weight compounds (e.g., lactate, acetate, methanol, or glucose) present in natural organic matter, fuel hydrocarbons, or less-chlorinated solvents (as discussed below).

#### 4.2.3.2 Electron Donor Reactions

Under aerobic conditions some chlorinated solvents can be utilized as the primary substrate (i.e., electron donor) in biologically mediated redox reactions (McCarty and Semprini, 1994). In this type of reaction, the facilitating microorganism obtains energy and organic carbon from the degraded compound. In contrast to reactions in which the chlorinated compound is used as an electron acceptor, only the least oxidized chlorinated solvents (e.g., VC, DCE, or chlorobenzene) may be utilized as electron donors in biologically mediated redox reactions. Also, chlorinated benzenes with up to four chlorines (i.e., CB, DCB isomers, TCB isomers, and TeCB isomers) have been shown to act as electron donors under aerobic conditions.

For example, while Murray and Richardson (1993) write that microorganisms are generally believed to be incapable of growth using TCE and PCE, other less chlorinated CAHs have been shown to be used as substrates. Davis and Carpenter (1990) describe the aerobic oxidation of VC in groundwater. McCarty and Semprini (1994) describe investigations in which VC and 1,2-DCA were shown to serve as primary substrates. These authors also document that dichloromethane has the potential to function as a primary substrate under either aerobic or anaerobic environments. Klier et al. (1996) describe aerobic mineralization of all three isomers of DCE. In addition, Bradley and Chapelle (1996) show evidence of oxidation of VC under iron-reducing conditions so long as there is sufficient bioavailable iron(III). CAH oxidation may be characterized by a loss of solvent mass, a decreasing molar ratio of daughter solvents to source solvent compounds, and rarely, the presence of chloromethane.

CB and polychlorinated benzenes (up to and including TeCB) have been shown to be biodegradable under aerobic conditions. Several studies have shown that bacteria are able to utilize CB (Reineke and Knackmuss, 1984), 1,4-DCB (Reineke and Knackmuss, 1984; Schraa et al., 1986; Spain and Nishino, 1987), 1,3-DCB (de Bont et al., 1986), 1,2-DCB (Haigler et al., 1988), 1,2,4-TCB (van der Meer et al., 1987; Sander et al., 1991), and 1,2,4,5-TeCB (Sander et al., 1991) as primary growth substrates in aerobic systems. Nishino et al. (1994) note that aerobic bacteria able to grow on chlorobenzene have been detected at a variety of chlorobenzene-contaminated sites, but not at uncontaminated sites. Spain (1996) notes that this provides strong evidence that the bacteria are selected for their ability to derive carbon and energy from chlorobenzene degradation in situ. The pathways for all of these reactions are similar, and are also similar to that of benzene (Chapelle, 1993; Spain, 1996). The only significant difference between aerobic oxidation of chlorobenzenes and benzene is the required elimination of chlorine. Anaerobic oxidation of chlorobenzenes has not been

conclusively documented, although little work has been done on the subject (Spain, 1997).

#### 4.2.3.3 Cometabolism

When a chlorinated solvent is biodegraded through cometabolism, it serves as neither an electron acceptor nor a primary substrate in a biologically mediated redox reaction. Instead, the degradation of the compound is catalyzed by an enzyme or cofactor that is fortuitously produced by organisms for other purposes. The organism receives no known benefit from the degradation of the solvent; rather the cometabolic degradation of the solvent may in fact be harmful to the microorganism responsible for the production of the enzyme or cofactor (McCarty and Semprini, 1994).

Cometabolism is best documented for chlorinated ethenes in aerobic environments, although it potentially could occur with other chlorinated solvents or under anaerobic conditions. Aerobic degradation pathways for chlorinated ethenes are illustrated on Figure 4.3. It has been reported that under aerobic conditions chlorinated ethenes, with the exception of PCE, are susceptible to cometabolic degradation (Murray and Richardson, 1993; Vogel, 1994; McCarty and Semprini, 1994). Vogel (1994) further elaborates that the cometabolism rate increases as the degree of halogenation decreases.

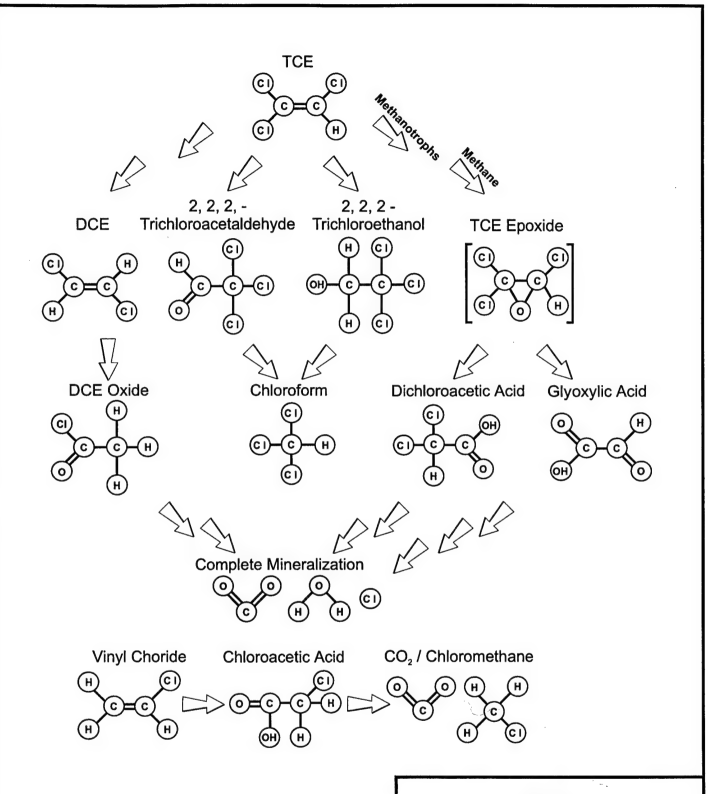
In the cometabolic process, TCE is indirectly transformed by bacteria as they use BTEX or another substrate to meet their energy requirements. Therefore, TCE does not enhance the degradation of BTEX or other carbon sources, nor will its cometabolism interfere with the use of electron acceptors involved in the oxidation of those carbon sources. Given this relationship, it would follow that depletion of suitable substrates (BTEX or other organic carbon sources) likely limits cometabolism of CAHs.

#### 4.2.3.4 Behavior of Chlorinated Solvent Plumes

Chlorinated solvent plumes can exhibit three types of behavior depending on the amount of solvent, the amount of organic (native and/or anthropogenic) carbon in the aquifer, the distribution and concentration of natural electron acceptors, and the types of electron acceptors being utilized. Individual plumes may exhibit all three types of behavior in different portions of the plume. The different types of plume behavior are summarized below.

#### 4.2.3.4.1 Type 1 Behavior

Type 1 behavior occurs where the primary substrate is anthropogenic carbon (e.g., BTEX or landfill leachate), and this anthropogenic carbon drives reductive dechlorination. When evaluating natural attenuation of a plume exhibiting type 1 behavior the following questions must be answered:



#### FIGURE 4.3

## AEROBIC DEGRADATION

OU-4 RNA TS Shaw AFB, South Carolina

PARSONS ENGINEERING SCIENCE, INC.

Denver, Colorado

- 1) Is the electron donor supply adequate to allow microbial reduction of the chlorinated organic compounds? In other words, will the microorganisms "strangle" before they "starve" [i.e., will they run out of chlorinated aliphatic hydrocarbons (electron acceptors) before they run out of primary substrate (anthropogenic carbon)]?
- 2) What is the role of competing electron acceptors (e.g., DO, nitrate, iron(III) and sulfate)?
- 3) Are VC and other less-chlorinated solvents being oxidized, or are they being reduced?

Type 1 behavior results in the rapid and extensive degradation of the highly chlorinated solvents such as PCE, TCE, or polychlorinated benzenes.

#### 4.2.3.4.2 Type 2 Behavior

Type 2 behavior dominates in areas that are characterized by relatively high concentrations of biologically available native organic carbon. This natural carbon source drives reductive dehalogenation (i.e., the primary substrate for microorganism growth is native organic carbon). When evaluating natural attenuation of a Type 2 chlorinated solvent plume, the same questions as those posed in the description of Type 1 behavior must be answered. Type 2 behavior generally results in slower biodegradation of the highly chlorinated solvents than Type 1 behavior, but under the right conditions (e.g., areas with high natural organic carbon contents), this type of behavior also can result in rapid degradation of these compounds.

#### 4.2.3.4.3 Type 3 Behavior

Type 3 behavior dominates in areas that are characterized by low concentrations of native and/or anthropogenic carbon, and concentrations of DO that are greater than 1.0 milligram per liter (mg/L). Under these aerobic conditions reductive dehalogenation will not occur. Thus there is little or no removal of PCE and TCE. Biodegradation may proceed via the much slower process of cometabolism, but will be limited by the low concentrations of native or anthropogenic carbon. The most significant natural attenuation mechanisms for CAHs will be advection, dispersion, and sorption. However, VC can be rapidly oxidized under these conditions; DCE may be oxidized; CBs may be oxidized; and cometabolism also may occur.

#### 4.2.3.4.4 Mixed Behavior

As mentioned above, a single chlorinated solvent plume can exhibit all three types of behavior in different portions of the plume. This can be beneficial for natural biodegradation of chlorinated solvent plumes. For example, Wiedemeier *et al.* (1996a) describe a CAH plume at Plattsburgh AFB, New York that exhibits Type 1 behavior in the source area and Type 3 behavior downgradient from the source. The best scenario involves a plume in which PCE, TCE, and DCE are reductively dehalogenated (Type 1 or Type 2 behavior), then VC is oxidized (Type 3 behavior), either aerobically or via iron reduction. VC is oxidized to carbon dioxide in this type of plume and does not

accumulate. The following sequence of reactions occurs in a plume that exhibits this type of mixed behavior:

$$PCE \rightarrow TCE \rightarrow DCE \rightarrow VC \rightarrow Carbon Dioxide$$

In general, the TCE, DCE, and VC may attenuate at approximately the same rate, and thus these reactions may be confused with simple dilution. Note that no ethene is produced during this reaction. VC is removed from the system much faster under these conditions than it is under VC-reducing conditions.

A less desirable scenario, but one in which all contaminants may be entirely biodegraded, involves a plume in which all CAHs are reductively dehalogenated via Type 1 or Type 2 behavior. VC is reduced to ethene, which may be further reduced to ethane or methane. The following sequence of reactions occur in this type of plume:

$$PCE \rightarrow TCE \rightarrow DCE \rightarrow VC \rightarrow Ethene or Ethane$$

This sequence has been investigated by Freedman and Gossett (1989). In this type of plume, VC degrades more slowly than TCE, and thus tends to accumulate.

#### 4.2.4 Abiotic Degradation of Chlorinated Solvents

Chlorinated solvents dissolved in groundwater may also be degraded by abiotic mechanisms, although the reactions may not be complete and often result in the formation of a toxic intermediate. The most common abiotic reactions affecting chlorinated solvents are hydrolysis and dehydrohalogenation. Hydrolysis is a substitution reaction in which a halogen substituent is replaced with a hydroxyl (OH) group from a water molecule. Dehydrohalogenation is an elimination reaction in which a halogen is removed from a carbon atom, followed by removal of a hydrogen atom from an adjacent carbon atom, with a double bond between the carbon atoms being produced. Other possible reactions include oxidation and reduction, although Butler and Barker (1996) note that no abiotic oxidation reactions involving common halogenated solvents have been reported in the literature. They also note that reduction reactions are most commonly microbially mediated.

Hydrolysis of chlorinated methanes and ethanes has been well-demonstrated in the literature (e.g., Vogel et al., 1987; Jeffers et al., 1989; Vogel, 1994; Butler and Barker, 1996). The likelihood that a solvent will hydrolyze is partly dependent upon the number of halogen substituents, typically with fewer halogens resulting in more rapid hydrolysis. Dehydrohalogenation, on the other hand, is more likely to take place as the number of halogen substituents increases. One common solvent for which abiotic degradation reactions have been well-documented is 1,1,1-TCA. 1,1,1-TCA may be transformed through a series of abiotic processes (including hydrolysis) to acetic acid. Vogel and McCarty (1987) demonstrated that 1,1,1-TCA can also dehydrohalogenate to 1,1-DCE, and Jeffers et al. (1989) demonstrated that 1,1,2-TCA also reacts in the same manner to form 1,1-DCE. Also, once TCA is reductively dehalogenated to chloroethane, it can then hydrolyze to ethanol (Vogel and McCarty, 1987).

Butler and Barker (1996) note that attributing changes in the presence, absence, or concentration of halogenated solvents to abiotic processes is usually difficult, particularly on the field scale. Solvents may undergo both biotic and abiotic degradation, and discerning the effects of each mechanism (at the field scale), if possible, would be very difficult. Also, as Butler and Barker (1996) note, the breakdown products of some reactions such as hydrolysis (e.g., acids and alcohols) may be more easily degraded (biotically or abiotically), resulting in limited accumulation; these potentially transient products also require additional analyses that may not be feasible for a field investigation. This makes field evidence to demonstrate hydrolysis very difficult to collect and interpret, and Butler and Barker (1996) note that such evidence has not been successfully collected and presented. Evidence of dehydrohalogenation is also difficult to collect, although the presence of 1,1-DCE in conjunction with 1,1,1-TCA can provide a tentative indication that the process is ongoing, and strong evidence where it is known that no DCE was released.

#### 4.3 DISTRIBUTION OF HYDROCARBONS AND DAUGHTER PRODUCTS

The first step for evaluating the occurrence and methods of biodegradation of chlorinated hydrocarbons and fuel hydrocarbons is to look at the distribution of target compounds and the products of biodegradation of those compounds. At many sites such as OU-4, dissolved groundwater contamination consists of a mixture of compounds, some of which may be electron donors, some of which may be electron acceptors, and some of which may be either donors or acceptors. It is therefore important to evaluate the distribution of all relevant compounds and the spatial relationships between those plumes.

For example, it is useful to determine if a source of dissolved electron donors (e.g., BTEX) coincides with a plume of dissolved contaminants that are most likely to be used as electron acceptors. The donors are needed to fuel biodegradation reactions that will either use some contaminants as electron acceptors or that will produce geochemical conditions that make the use of contaminants as electron acceptors more favorable. Also, because reductive dehalogenation is the most common biodegradation reaction involving chlorinated hydrocarbons, it is useful to look for the presence of compounds that are most likely the product of biodegradation reactions, such as *cis*-1,2-DCE or VC. These compounds are rarely released to groundwater, but are products of reductive dehalogenation. In addition, if *cis*-1,2-DCE is clearly the most common isomer of DCE, then the occurrence of reductive dehalogenation can be inferred.

The following sections present contaminant and daughter product distributions at OU-4, providing the initial evidence that dissolved fuels and chlorinated solvents are biodegrading. This evidence alone is not conclusive, but when combined with geochemical evidence presented in later sections, it is a significant indicator of biodegradation.

#### 4.3.1 Fuel Hydrocarbons

Fuel hydrocarbons detected in groundwater at Site OU-4 include all of the BTEX compounds, as well as TMB isomers and fuel carbon. Results for fuel hydrocarbon analyses performed on samples collected in May 1997 are presented in Table 4.3, and the distribution of total BTEX is shown on Figure 4.4. Where detected, total BTEX

# GROUNDWATER QUALITY DATA SUMMARY FOR BTEX, TMBs, AND FUEL CARBON

OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

				Ethyl-	Total	Total	1,3,5-	1,2,4-	1,2,3-	Fuel
	Date	Benzene	Toluene	Benzene	Xylenes	BTEX	TMB	TMB	TMB	Carbon
Sample ID	Sampled	$(\mu g/L)^{a'}$	(µg/L)	(µg/L)	(µg/L)	(µg/L)	(µg/L)	$(\mu g/L)$	$(\mu g/\Gamma)$	$(\mu g/L)$
MW-105	5/15/97	7.07	7.1	37.9	224.9	340.6	3.5	137	52.7	930
MW-106	5/14/97	< 1 <sub>b</sub> /	<	5.6	4.1	8.7	$ND_{o}$	2.8	N Q	14.0
MW-107	5/14/97	N N	R	N Q	N N	R	N N	QN	Q.	N Q
MW-111	5/15/97	R	N N	QN Q	QN Q	R	N N	N Q	R	N Q
MW-112	2/16/97	229	438	495	1737	2899	177	521	193	2080
MW-112A	2/16/97	R	R	QN.	QQ.	R	R	QN	R	ND
MW-113	5/16/97	<u>R</u>	R	<u>^</u>	1.9	2.9	R	QN.	N N	<u>^</u>
MW-114	5/15/97	6.3	12.2	8.99	219.9	295.2	14.3	74.9	20.3	623
MW-115	5/15/97	157	393	113	588	1251	26.0	161	20.2	2350
MW-115A	5/15/97	N N	N N	Q.	QN Q	Q.	R	S	S	NO.
MW-116	5/14/97	QN Q	R	QN	N	N Q	<u>R</u>	Ω	R	N Q
MW-116A	5/14/97	Q.	R	ND	QN	N Q	R	S	R	ND
MW-117	5/15/97	30.0	89.5	72.8	396.5	588.8	13.4	90.5	41.1	1290
MW-118	5/15/97	Q.	R	N Q	QN	Q.	Q.	S	R	ND
TW-101	5/14/97	QN	N N	Q	N Q	N N	R	Q.	Q Q	QN
TW-120	5/11/97	Ð	2.1	S	<u>R</u>	2.1	R	Q.	N Q	1.8
TW-121	5/17/97	5.3	1.2	-	4	11.5	R	6.0	1.2	16.1
MPA	5/16/97	14.5	943	403	1934	3294.5	152	321	506	5540
MPB	2/16/97	7.1	917	224	1101	2249.1	118	326	144	4130
MPC	5/16/97	81.7	1685	242	1279	3287.7	118	328	148	4730
MPD	5/16/97	19.3	159	142	496	816.3	40.4	204	44.9	1720
MPE	5/11/97	QN	R	R	2.0	2.0	R	QN.	R	< <del>1</del> >
TMP-1	5/11/97	R	£	Q.	R	Ð	£	QN N	Q	ND
TMP-2	5/11/97	12.5	2.3	2.4	4.7	21.9	6.0	4.2	6.7	127
TMP-3	5/11/97	12.4	<u>R</u>	R	2.2	14.6	R	N N	4.5	60.4
TMP-4	2/16/97	10.1	<u>~</u>	8.4	8.5	28.0	<u>~</u>	13.2	<u>~</u>	207
TMP-5	5/16/97	28.2	6.6	77.2	128.3	243.6	19	72.3	27.7	529
SWS-1	5/15/97	S	QN Q	S S	N N	Q.	N Q	S	Q Q	ND
SWS-2	5/15/97	Q	Q.	N Q	Q	QN	QN	S	N Q	ND
SWS-3	5/15/97	Q	Q.	N	QN	Q	QN	N	QN	ND
SWS-4	5/15/97	Q	QN.	R	ND	QN	N N	ND	N	ND
SWS-5	5/15/97	QN	Ð	ND	ND	R	QN ON	N N	ND	ND
1/2/4	1:40=									

 $<sup>^{</sup>a\prime}$  µg/L = micrograms/liter.  $^{b\prime}$  <1 = Below limit of quantitation, 1 µg/L.

o' ND = Analyte not detected.

concentrations ranged from 2.0 to 3,295 micrograms per liter ( $\mu$ g/L). Detected benzene concentrations ranged from <1 to 229  $\mu$ g/L, with concentrations exceeding the USEPA (1996) maximum contaminant level (MCL) of 5  $\mu$ g/L at 14 locations. Toluene concentrations ranged from <1 to 1,685  $\mu$ g/L and exceeded the MCL of 1,000  $\mu$ g/L at 1 location. Ethylbenzene (ranging from <1 to 403  $\mu$ g/L) and xylenes (ranging from 2 to 1,934  $\mu$ g/L) did not exceed MCLs at the site (Table 4.3).

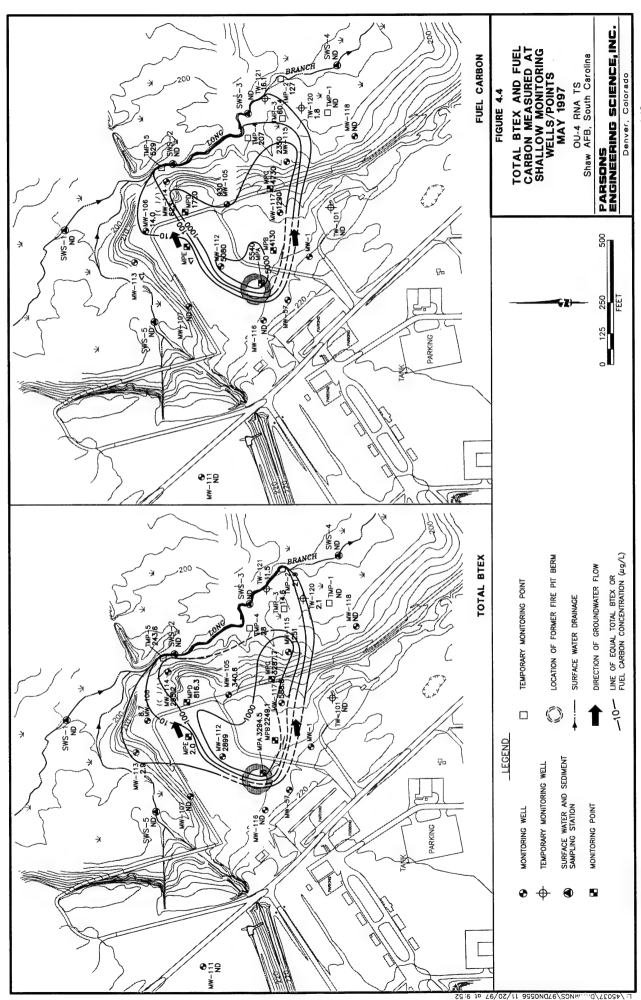
The groundwater BTEX plume is double-lobed with one lobe extending to the east/southeast and the other to the northeast. Figure 4.4 shows that groundwater BTEX concentrations are highest along the east/southeast plume lobe. Monitoring points MPA and MPC in this lobe have the two highest BTEX concentrations detected at OU-4, which coincide with two known contaminant sources at the site (MPA is in the former bermed burn pit, and MPC contained mobile LNAPL). The groundwater BTEX concentrations in the northeast plume lobe steadily decline from the fire training pit to the Long Branch Creek and suggest a single groundwater source near the fire training pit. Figure 4.5 breaks the total BTEX plume down into the individual BTEX The maximum benzene, ethylbenzene, and xylenes compounds were compounds. located along the northeastern plume lobe, whereas the maximum toluene concentration was located along the east/southeast plume lobe. Individual TMB isomers also were detected at 13 wells/points, with concentrations ranging from < 1 µg/L to 328 µg/L (Table 4.3). BTEX were detected in all samples containing TMB.

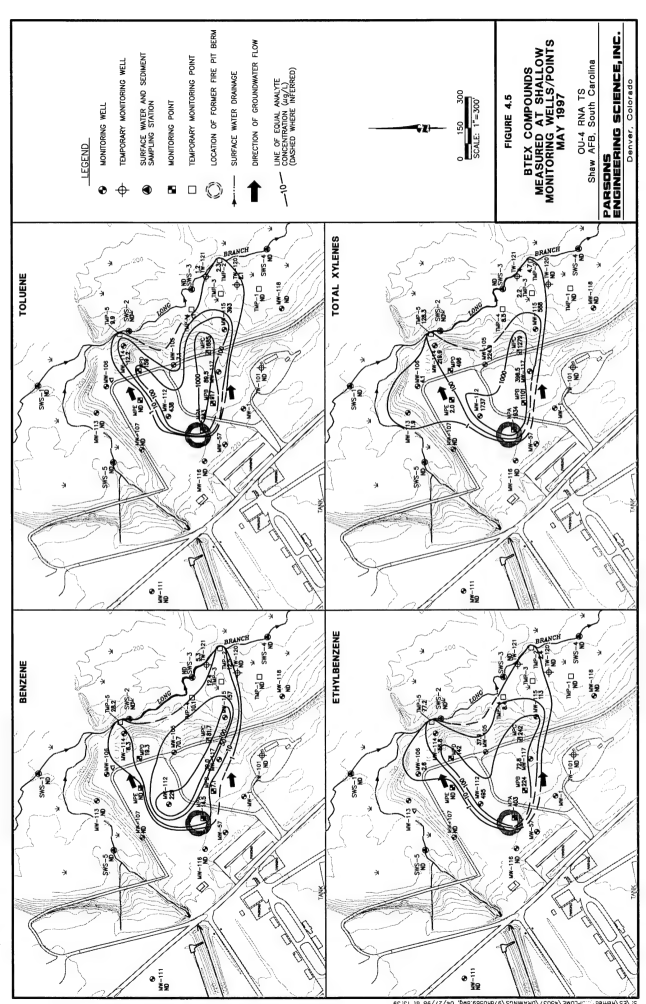
The distribution of total fuel carbon (normalized to JP-4), is shown on Figure 4.4, represents the mass of fuel carbon dissolved in groundwater (fuel carbon mass was divided by approximately 0.85 to estimate TPH mass). The fuel carbon plume is similar in size and shape to the total BTEX plume. The average ratio between fuel carbon concentrations and BTEX compounds is approximately 2 to 1, indicating that BTEX comprises nearly 50 percent of the total dissolved fuel hydrocarbon concentration. Plausible fractions of BTEX dissolving from a fresh JP-4 source residing on water range from 41 to 82 percent of the total dissolved fuel hydrocarbon (Smith et al., 1981).

#### 4.3.2 Chlorinated Ethenes

Chlorinated ethenes detected in groundwater at OU-4 include PCE, TCE, DCE isomers, and VC. Data for chlorinated ethenes and other chlorinated compounds are presented in Table 4.4. May 1997 distributions of PCE, TCE, cis-1,2-DCE, and VC are shown on Figure 4.6.

Groundwater PCE concentrations measured in May 1997 ranged from <1 to 90.7  $\mu$ g/L and exceeded the USEPA (1996) MCL of 5  $\mu$ g/L at two locations. The source of PCE appears to be centered around monitoring point MPC where a fuel/solvent mobile LNAPL was observed (Section 4.1.1). PCE was not detected at monitoring point MPA located within the fire training pit. Temporary monitoring points located downgradient and near Long Branch Creek had mostly non-detectable PCE concentrations, with the exception of monitoring point TMP-2, which had a low PCE detection below the quantitation limit. PCE does not appear to be discharging to Long Branch Creek, nor was it detected at the three deep wells screened in the surficial aquifer at the site (MW-112A, -115A, and -116A). Therefore, the potential for significant receptor exposure to PCE appears to be low.





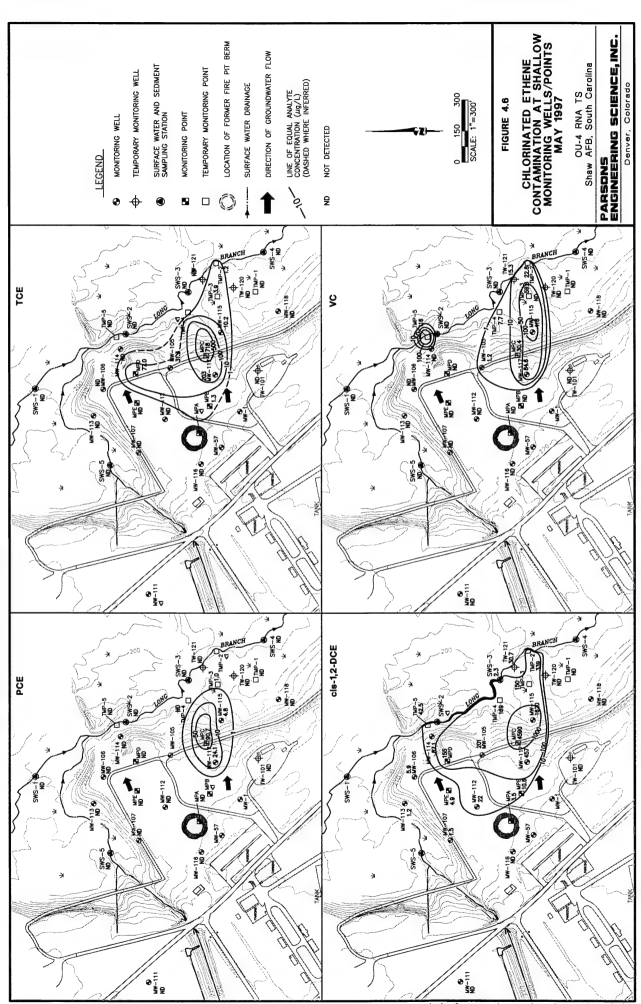
# GROUNDWATER AND SURFACE WATER QUALITY DATA SUMMARY FOR CHLORINATED ETHENES, ETHANES, AND CHLOROFORM

SHAW AFB, SOUTH CAROLINA **OU-4 RNA TS** 

										,	
				cis-	trans-	1,1-	Vinyl	1,1,1-	1,1-	1,2-	
Sample	Date	PCE	TCE	1,2-DCE	1,2-DCE	DCE	Chloride	TCA	DCA	DCA	Chloroform
Ð	Sampled	$(\mu g/L)^{u'}$	$(\mu g/L)$	(µg/L)	$(\mu g/L)$	(µg/L)	$(\mu g/L)$	(µg/L)	$(\mu g/L)$	(µg/L)	(µg/L)
MW-105	5/15/97	< 1 <sub>b</sub> /	31.3	201	ND/c	< 1	1.2	ND	<1	ND	47.9
MW-106	5/14/97	QN QN	NO	5.9	ON	<u>~</u>	QN Q	N ON	3.0	Q.	R
MW-107	5/14/97	QN	QN ON	1.5	ND	N Q	NO	QN Q	1.1	QN Q	Q.
MW-111	5/15/97	QN	1 >	S	ND	QN	S S	N Q	N Q	Q	QN
MW-112	2/16/97	Q.	NO NO	22.0	ND	2.2	QN Q	\ \ 	64.8	QN	2.3
MW-112A	5/16/97	S	QN.	Q.	ON	S S	QN ON	QN	ND	QN	ND
MW-113	5/16/97	QN ON	ND	1.2	ND	ND	ON ON	QN Q	<1	Q	N N
MW-114	5/15/97	ND	QX	87.1	NO	2.1	QZ	NO	58.8	QN	<u>^</u>
MW-115	5/15/97	4.8	10.2	1620	N ON	368	416	2030	8700 <sub>4</sub> /	QN	<u>~</u>
MW-115A	5/15/97	N ON	× 1	QN ON	ON	2.1	ON O	ND	<1	QN	ND
MW-116	5/14/97	ND	Q	Q.	ND	QN QN	N ON	ND	ND	QN ON	ND
MW-116A	5/14/97	QN	S	N QN	ND	N ON	ND	ND	ND	N Q	ND
MW-117	5/15/97	24.1	203	457	ND	148	84.6	4690 <sup>d/</sup>	481	S S	<1
MW-118	5/15/97	ND	N Q	Q.	N ON	Q Q	QN.	<u>^</u>	Q.	QN ON	ND ND
TW-101	5/14/97	NO	ND	Ð	ND ON	NO	N	N Q	QN	QZ	ND
TW-120	5/17/97	QX	N	ON N	ND	ND	N QN	ND QN	QN ON	ND	ND
TW-121	5/17/97	N ON	N Q	30.7	2.1	7.3	15.3	N Q	1.5	ND	R
MPA	5/16/97	QN ON	<u> </u>	3.5	N Q	4.3	NO	8.4	30.8	N Q	7
MPB	2/16/97	<u>~</u>	1.3	10.6	S	4.7	QN	411	15.4	N N	<1
MPC	5/16/97	90.7	718	4590 <sup>4/</sup>	ND	748	30.4	$23000^{d/}$	1060	ND	1.2
MPD	5/16/97	QN	77.0	155	NO	1.7	N N	1.3	53.3	NO NO	<u>^</u>
MPE	5/17/97	ON	QN QN	4.9	QN	QN	QN ON	<1	1.4	S S	ND
TMP-1	5/17/97	NO	ND	S	ND	QN QN	ND	ND	QN Q	QZ QZ	ND ND
TMP-2	5/17/97	<u>^</u>	1.2	339	ND	116	22.8	78.6	1070	NO	ND
TMP-3	5/17/97	1.0	3.9	150	N Q	40.2	59.6	60.5	1153	2.1	QN Q
TMP-4	5/16/97	N Q	<1	169	ND	3.5	7.7	<1	64.9	Z	NO ON
TMP-5	2/16/97	Q.	QN Q	42.5	N Q	1.8	116	Q.	3.5	QN	QN QN
SWS-1	5/15/97	QN ON	QN ON	ND	QN ON	ND	N Q	QN	S	QN	QN Q
SWS-2	5/15/97	ND	ND	\ \ !	NO	ND	N	QN	ND QN	NO	N Q
SWS-3	5/15/97	QN	ND	2.3	ND	\ 1	N Q	< 1 ×	1.0	ΩN	Q.
SWS-4	5/15/97	N ON	QN QN	1 > 1	ND	N N	ND	ND	\ \ 1	ND	QN
SWS-5	5/15/97	ND	QN	R	ND QN	QN.	ND	ND	Q.	QN	ND
a/ ug/L = Micro	ug/L = Micrograms per Liter.		o' ND = Analy	nalyte not detected							

ND = Analyte not detected.

 $<sup>^{</sup>d'}$  Analyte values reported based on a correction for 1/10 dilution.  $^{\prime\prime\prime}$  µg/L = Micrograms per Liter.  $^{\prime\prime}$  <1 = Below limit of quantitation, 1 µg/L.



Groundwater TCE concentrations were slightly more widespread than PCE concentrations (Figure 4.6) and are likely the product of more than one source. The primary TCE source appears to be centered around monitoring point MPC where, like PCE, the maximum TCE concentration at the site was measured (718  $\mu$ g/L). The reductive dechlorination of PCE likely contributed to the elevated TCE concentration at this location; however, TCE likely was used as a combustible solvent during fire training exercises. A second possible TCE source is located at or upgradient from monitoring point MPD. A product sheen was detected at MPD during groundwater sampling in May 1997 (Section 4.1.1). TCE does not appear to be a discharge threat to Long Branch Creek because the single highest TCE concentration measured near the creek was 1.2  $\mu$ g/L at monitoring point TMP-2. TCE exceeded the USEPA (1996) MCL of 5  $\mu$ g/L at five sampling locations.

The total DCE plume spanned a slightly larger area than the PCE and TCE plumes (Figure 4.6). Low DCE concentrations were measured in groundwater as far north as monitoring well MW-113 and suggest the former presence of low-level PCE and/or TCE contamination in the shallow aquifer north of the fire-training pit. The maximum DCE concentration was detected at point MPC (5,338 µg/L), where PCE and TCE contamination was also greatest. Cis-1,2-DCE ranged from 45 to 100 percent of the total DCE concentration (average of 85 percent). Figure 4.6 shows cis-1,2-DCE concentrations, which represents the majority of the DCE plume mass. concentration ranges for the other DCE isomers were 2.1 µg/L for trans-1,2-DCE (measured at well TW-121 only) and <1 to 368 μg/L for 1,1-DCE. Cis-1,2-DCE exceeded the USEPA (1996) MCL of 70 µg/L at nine locations. 1,1-DCE exceeded the its MCL of 7 µg/L at five locations, and trans-1,2-DCE concentrations did not exceed its MCL. A high proportion of cis-1,2-DCE relative to trans-1,2-DCE is expected where TCE is reductively dehalogenated to DCE. A fraction of trans-1,2-DCE comparable to or exceeding cis-1,2-DCE would suggest the possibility of DCE as a source solvent because the trans isomer is more common when DCE is used for commercial applications. The relative abundance of cis-1,2-DCE is therefore a good indication that the initial reductive dehalogenation step of TCE is taking place in groundwater at Site OU-4 (Bouwer, 1994). 1.1-DCE also may result from the reductive dehalogenation of TCE, but the 1,1-DCE plume exceeding 10 µg/L coincides with and is slightly downgradient from the 1,1,1-TCA plume exceeding 10 μg/L (Section 4.3.3). Furthermore, these plumes are entirely within the east/southeast lobe of the plume, whereas elevated TCE concentrations also are present in the northeast lobe. This suggests that most of the 1,1-DCE results from dehydrohalogenation of 1,1,1-TCA.

The distribution of VC in site groundwater is shown on Figure 4.6. VC concentrations at the site ranged from 1.2 to 416  $\mu$ g/L (Table 4.4). Concentrations exceeded the MCL of 2  $\mu$ g/L at eight of nine locations where detected. The maximum VC concentration was detected at monitoring well MW-115, which is approximately 80 feet from the maximum concentrations of PCE, TCE, and DCE, detected at monitoring point MPC. The presence of VC in groundwater is also a good indicator of the occurrence of reductive dehalogenation, because VC is typically a gas at most working temperatures [the boiling point of VC is about 7°F (Montgomery, 1996)]. Because of this, VC is not commonly used as a solvent, but instead is used for plastic manufacturing. In addition, the low boiling point indicates that when liquid VC is

spilled, most of the mass is likely to volatilize before reaching groundwater unless there is a large-volume release.

#### 4.3.3 Chlorinated Ethanes

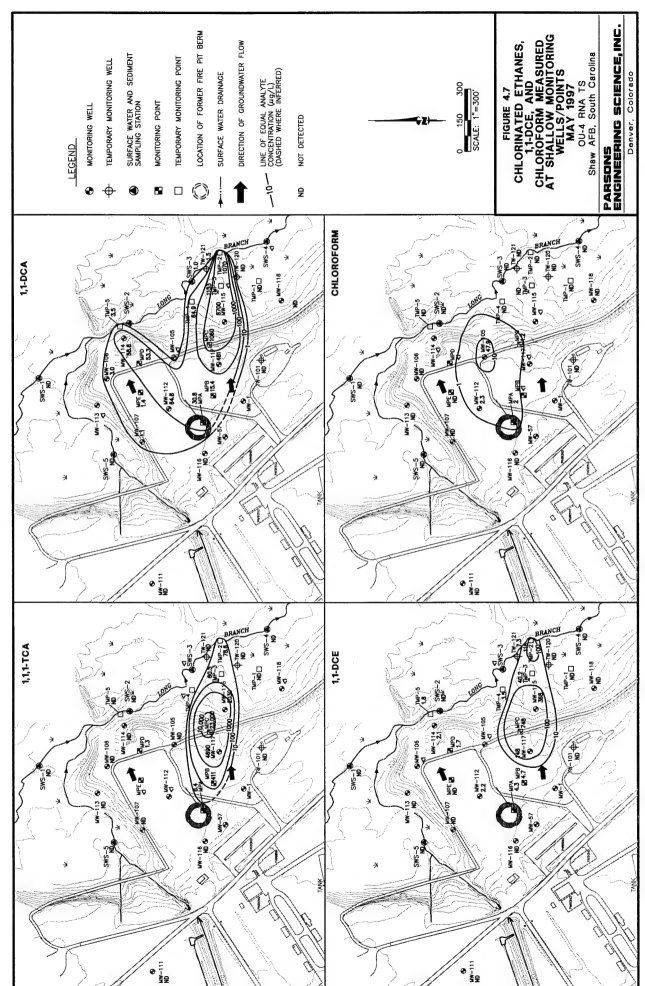
Chlorinated ethanes detected in groundwater at OU-4 include 1,1,1-TCA, 1,2-DCA, and 1,1-DCA (Table 4.4). The distributions of these compounds are shown on Figure 4.7. Relatively high concentrations of 1,1,1-TCA and 1,1-DCA are present at the site, with the maximum concentrations of both compounds measured at monitoring point MPC or monitoring well MW-115. The concentration ranges of detected chlorinated ethane compounds are <1 to 23,000  $\mu$ g/L for 1,1,1-TCA, <1 to 8,700  $\mu$ g/L for 1,1-DCA, and a single detection of 2.1  $\mu$ g/L for 1,2-DCA. The USEPA (1996) MCL of 200  $\mu$ g/L for 1,1,1-TCA was exceeded at three locations. MCLs are not available for the DCA isomers. In general, chlorinated ethanes were detected in the same wells as the chlorinated ethenes.

The 1,1,1-TCA plume was centered around monitoring point MPC where a mixture of mobile LNAPL containing fuel hydrocarbons and chlorinated solvent was observed during May 1997 field work. The mobile LNAPL source was not analyzed for TCA isomers, but is the most likely source of contamination. The presence of both 1,1-DCA and fuel hydrocarbons at significant concentrations suggests that the reductive dehalogenation of 1,1,1-TCA is occurring. Additionally, 1,1-DCA is not commonly used as a solvent, whereas 1,2-DCA is; therefore, the presence of 1,1-DCA rather than 1,2-DCA (nearly absent from the site) further suggests that reductive dechlorination of 1,1,1-TCA is ongoing, particularly in the east/southeast plume lobe. The presence of 1,1-DCE was generally observed at every location where 1,1,1-TCA was detected. 1,1-DCE is the least common daughter product generated as a result of TCE dechlorination, whereas cis-1,2-DCE followed by trans-1,2-DCE are typically more commonly observed daughter products. The widespread occurrence of cis-1,2-DCE and absence of trans-1,2-DCE suggests that 1,1-DCE was not generated through the reductive dechlorination of TCE. Rather, 1,1-DCE has probably formed through the dehydrohalogenation of 1,1,1-TCA at locations where both are present. The abiotic reaction is relatively slow compared to reductive dehalogenation, which could explain why more 1,1-DCA (product of 1,1,1-TCA reductive dechlorination) than 1,1-DCE (product of 1,1,1-TCA dehydrohalogenation) was observed.

#### 4.3.4 Chlorobenzenes

Chlorobenzenes detected in groundwater at Site OU-4 include 1,2-DCB, 1,3-DCB, 1,4-DCB, and CB (Table 4.5). These compounds generally were detected in the same wells as the other CAHs, and plumes for all four compounds appear to originate from the vicinity of MPC and migrate exclusively in the east/southeast plume lobe (Figure 4.8). The chlorobenzenes ranged in concentration from 1.7 to 117  $\mu$ g/L for 1,2-DCB, <1 to 68.2  $\mu$ g/L for 1,3-DCB, <1 to 14.1 for 1,4-DCB, and <1 to 9.2  $\mu$ g/L for CB. The USEPA MCL for CB is 100  $\mu$ g/L, the MCL for 1,2-DCB and 1,3-DCB is 600  $\mu$ g/L, and the MCL for 1,4-DCB is 75  $\mu$ g/L (USEPA, 1991). None of the chlorobenzenes detected at OU-4 exceeded federal MCLs.

It is possible that some DCB is being reductively dechlorinated to CB in the vicinity of MPC where conditions may be the most reducing, although little information has

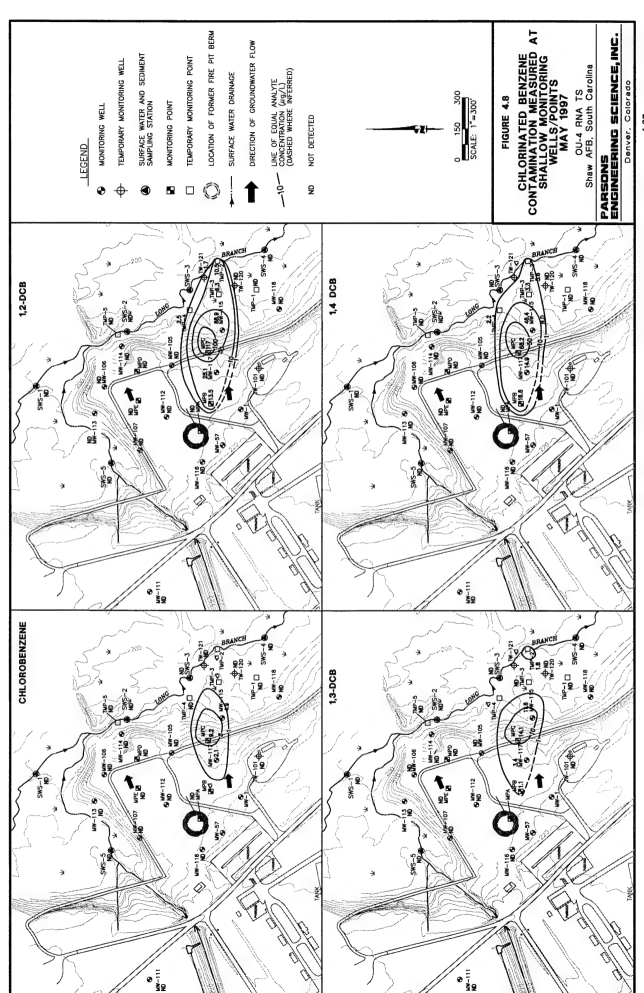


#### TABLE 4.5 GROUNDWATER QUALITY DATA SUMMARY FOR CHLORINATED BENZENES

		Chloro-	1,3-	1,4-	1,2-	Total Chlorinated
Sample	Date	benzene	DCB	DCB	DCB	Benzenes
ID	Sampled	(μg/L) <sup>a/</sup>	(μg/L)	(μg/L)	(μg/L)	(μg/L)
MW-105	5/15/97	$ND_{p}$	ND	ND	ND	0
MW-106	5/14/97	ND	ND	ND	ND	0
MW-107	5/14/97	ND	ND	ND	ND	0
MW-111	5/15/97	ND	ND	ND	ND	0
MW-112	5/16/97	ND	ND	ND	ND	0
MW-112A	5/16/97	ND	ND	ND	ND	0
MW-113	5/16/97	ND	ND	ND	ND	0
MW-114	5/15/97	ND	ND	ND	ND	0
MW-115	5/15/97	4.9	11.6	46.4	88.9	151.8
MW-115A	5/15/97	ND	ND	ND	ND	0
MW-116	5/14/97	ND	ND	ND	ND	0
MW-116A	5/14/97	ND	ND	ND	ND	0
MW-117	5/15/97	2.1	3.4	14.9	25.1	45.5
MW-118	5/15/97	ND	ND	ND	ND	0
TW-101	5/14/97	ND	ND	ND	ND	0
TW-120	5/17/97	ND	ND	ND	ND	0
TW-121	5/17/97	ND	< 1	< 1	1.7	1.7
MPA	5/16/97	ND	ND	ND	ND	0
MPB	5/16/97	< 1 <sup>c/</sup>	1.1	16.8	13.5	31.4
MPC	5/16/97	9.2	14.1	68.2	117	208.5
MPD	5/16/97	ND	ND	ND	ND	0
MPE	5/17/97	ND	ND	ND	ND	0
TMP-1	5/17/97	ND	ND	ND	ND	0
TMP-2	5/17/97	< 1	1.8	5.6	10.5	17.9
TMP-3	5/17/97	< 1	ND	3.3	6.3	9.6
TMP-4	5/16/97	ND	< 1	2.2	2.5	4.7
TMP-5	5/16/97	ND	ND	ND	ND	0
SWS-1	5/15/97	ND	ND	ND	ND	0
SWS-2	5/15/97	ND	ND	ND	ND	0
SWS-3	5/15/97	ND	ND	ND	ND	0
SWS-4	5/15/97	ND	ND	ND	ND	0
SWS-5	5/15/97	ND	ND	ND	ND	0

a/ μg/L = Micrograms per Liter.

b/ ND = Analyte not detected.



been published on this mechanism. As with chlorinated ethenes and ethanes, the degree of reductive dechlorination of chlorinated benzenes is expected to diminish with decreasing chlorination. Therefore, DCB and CB (least chlorinated of the chlorobenzenes) are least susceptible to reductive dechlorination. It is more likely that both CB and DCB reductions are being accomplished through oxidation reactions where the chlorobenzenes act as electron donors/substrates in a manner similar to BTEX biodegradation.

#### 4.3.5 Other Chlorinated Compounds

One other chlorinated compound was detected in groundwater at Site OU-4. Chloroform was detected at concentrations of <1 to 47.9 µg/L (Table 4.4 and Figure 4.7). None of the chloroform detections exceeded the USEPA (1996) MCL of 100 µg/L. Chloroform concentrations are generally located in the area between the two primary BTEX, chlorinated ethene, and chlorinated ethane plume lobes. These same locations have relatively lower BTEX and TCE concentrations, which is reflective of being located between the two primary contaminant plume lobes. Therefore, the aquifer may be more susceptible to oxygen enrichment from precipitation and the activity of aerobic microorganisms. Chloroform can result from the aerobic oxidation of TCE and TCE daughter products (e.g., 2,2,2-trichloracetaldehyde or 2,2,2trichloroethanol) produced during aerobic oxidation. Chloroform can also result from the reductive dechlorination of carbon tetrachloride (Criddle et al., 1990). Carbon tetrachloride was not detected at the site and is not suspected to be a source of chloroform contamination. Chloroform is used in industrial cleaning operations (e.g., for cleaning circuit boards, refrigerants, or as a solvent for natural products) and its use as a combustible waste is not documented at OU-4.

#### 4.3.6 Chloride

Chloride ions are removed from chlorinated solvents and enter solution during biodegradation, whether via reductive dehalogenation or aerobic oxidation. Therefore, chloride concentrations in groundwater should increase above background levels in areas where significant reductive dehalogenation or aerobic biodegradation is taking place. Chloride concentrations are presented in Table 4.6, and Figure 4.9 shows the May 1997 distribution of chloride in shallow groundwater at the site. Background chloride concentrations (measured in wells without contamination in shallow site groundwater) average approximately 3 mg/L. Chloride concentrations within the boundaries of the east/southeast chlorinated solvent plume lobe generally are elevated slightly above background, with a maximum concentration of 20.2 mg/L (at MW-115).

In conjunction with the distributions of CAH compounds shown on Figures 4.6 through 4.8, the elevated chloride concentrations observed at Site OU-4 are a good indication that CAHs are undergoing microbially mediated degradation. It is particularly noteworthy that the two highest chloride concentrations occur at and downgradient from MPC, where the highest CAH concentrations were detected.

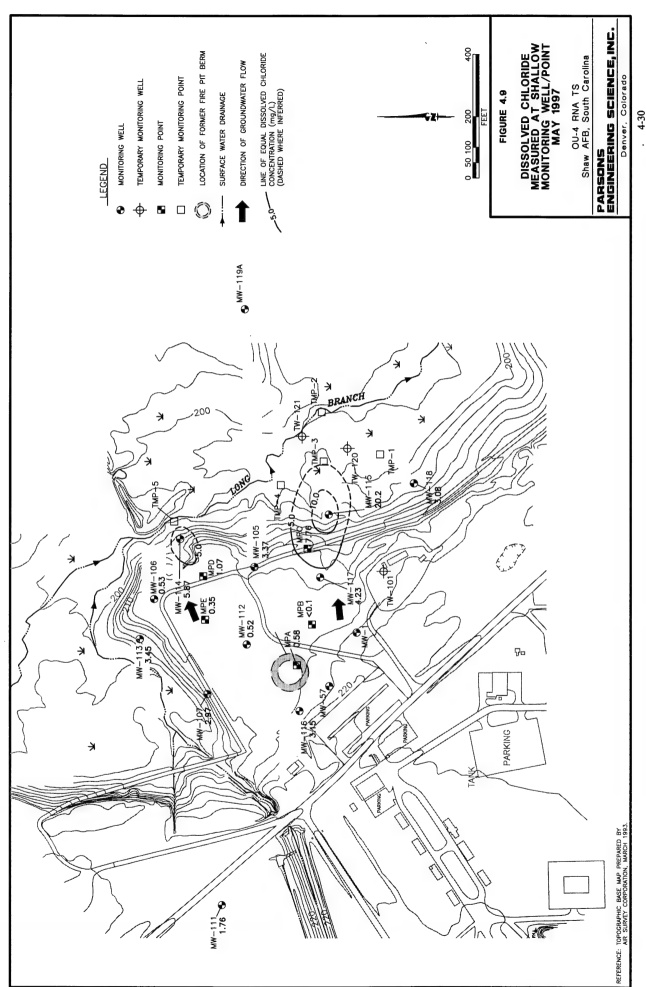
TABLE 4.6
GROUNDWATER ELECTRON ACCEPTORS/BYPRODUCTS

				A STITE	Ara, 500 tr	TOWN II	- T					
		Dissolved	Carbon			Nitrate +	Ferrous	Hydrogen				
Sample	Date	Oxygen	Dioxide	Chloride	Sulfate	Nitrite	Iron	Sulfide	Ammonia	Methane	Ethene	Ethane
ID	Sampled	$(mg/L)^{a'}$	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)
MW-105	5/15/97	0.2	88	3.37	5.57	< 0.05	3.08	< 0.1	2.18	3.59	NDb	N Q
MW-106	5/14/97	0.3	32	0.53	< 0.1	0.11	3.00	0.1	< 0.05	90.0	R	ND
MW-107	5/14/97	2.7	28	2.97	4.00	0.26	1.05	< 0.1	< 0.05	0.01	N N	ND ND
MW-111	5/15/97	8.0	56	1.76	< 0.1	0.59	Q	< 0.1	< 0.05	S	N N	R
MW-112	5/16/97	<0.1	120	0.52	< 0.1	< 0.05	7.00	0.5	0.83	6.05	R	ND
MW-112A	5/16/97	5.6	20	2.55	1.53	0.47	0.01	< 0.1	< 0.05	S	R	R
MW-113	5/16/97	3.8	22	3.45	< 0.1	0.28	0.13	< 0.1	< 0.05	0.00	N	ND
MW-114	. 5/15/97	0.4	36	5.87	2.75	< 0.05	2.15	0.5	< 0.05	1.14	N N	ND
MW-115	5/15/97	<0.1	84	20.2	4.30	< 0.05	5.90	0.5	1.52	2.46	0.01	0.00
MW-115A	5/15/97	7.4	22	3.92	< 0.1	0.99	0.08	< 0.1	< 0.05	N Q	R	R
MW-116	5/14/97	8.4	81	3.15	15.1	0.99	0.04	< 0.1	< 0.05	R	N	N
MW-116A	5/14/97	5.0	28	4.26	3.12	0.51	QN ON	< 0.1	< 0.05	N Q	R	R
MW-117	5/15/97	9.0	44	4.23	20.4	< 0.05	0.03	< 0.1	0.26	0.33	0.00	< 0.002
MW-118	5/15/97	2.5	34	3.08	60.6	0.79	ND	< 0.1	< 0.05	0.00	R	QN Q
TW-101	5/14/97	4.1	20	NAc	NA	1.20	0.84	< 0.1	0.78	0.15	ND	S
TW-120	5/17/97	0.3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
TW-121	5/17/97	3.1	40	NA	NA	< 0.05	4.00	< 0.1	6.35	4.54	N N	0.01
MPA	5/16/97	0.5	100	0.58	< 0.1	< 0.05	4.00	0.2	7.41	6.95	QN Q	QN
MPB	5/16/97	0.3	28	< 0.1	3.21	< 0.05	3.00	2	1.18	3.38	ND ND	N Q
MPC	5/16/97	0.5	70	7.76	< 0.1	< 0.05	4.00	0.3	0.43	0.07	< 0.003	0.01
MPD	5/16/97	0.5	NA	1.07	< 0.1	< 0.05	NA	NA	0.20	1.63	R	S
MPE	5/17/97	0.4	20	0.35	< 0.1	< 0.05	7.00	< 0.1	0.17	0.03	N N	QZ
TMP-1	5/17/97	3.0	30	NA	NA	NA	0.20	< 0.1	NA	0.00	R	R
TMP-2	5/17/97	0.2	80	NA	NA	< 0.05	8.20	0.3	0.39	0.34	< 0.003	R
TMP-3	5/17/97	0.2	40	NA	NA	0.41	0.18	< 0.1	< 0.05	0.19	< 0.003	Q.
TMP-4	2/16/97	<0.1	20	NA	NA	0.40	4.00	0.1	0.47	0.22	Q	S
TMP-5	2/16/97	0.1	20	NA	NA	< 0.05	4.00	< 0.1	0.37	1.71	< 0.003	N
a/ mo/I. = milliprams per liter.	ns per liter.		* Note: TV	V-101 was m	nislabeled as	MW-101 fe	or nitrate+ni	Note: TW-101 was mislabeled as MW-101 for nitrate+nitrite and ammonia.	nonia.			

Note: TW-101 was mislabeled as MW-101 for nitrate+nitrite and ammonia. mg/L = milligrams per liter.

 $<sup>^{</sup>b'}$  ND = Not detected.

 $<sup>\</sup>omega'$  NA = Not available.



#### 4.4 ADDITIONAL EVIDENCE OF BIODEGRADATION

As noted in Section 4.2, comparing concentrations and distributions of electron donors, electron acceptors, and byproducts of microbially mediated reactions can help indicate the types of processes operating at a site. In addition, other geochemical parameters, including ORP, alkalinity, and other changes in groundwater chemistry can provide supporting evidence. Volatile fatty acid concentrations also can be useful for interpreting and confirming biodegradation mechanisms.

#### 4.4.1 Electron Donors, Native Electron Acceptors, and Byproducts

When investigating the biodegradation of hydrocarbons, it is informative to evaluate the distribution of other compounds that are used in or produced by the microbially mediated reactions that facilitate contaminant degradation. The distribution of potential electron donors other than contaminants, such as dissolved native organic carbon, is useful for evaluating the feasibility of reductive dehalogenation or cometabolism (i.e., biodegradation reactions involving another substrate). Moreover, electron acceptor and metabolic byproduct data can clarify which processes may be facilitating contaminant degradation.

#### 4.4.1.1 Organic Carbon in Groundwater

Dissolved organic carbon can act as a source of electron donors during the reductive dehalogenation of CAHs. Dissolved TOC data from wells outside contaminant plumes can be used as an indicator of background levels of native carbon compounds.

TOC dissolved in groundwater was measured in some of the groundwater samples collected in May 1997. These concentrations are presented in Table 4.7 and shown on Figure 4.10. Dissolved TOC concentrations detected in shallow groundwater at OU-4 range from 1.05 to 144 mg/L. The highest concentration was measured at MPA, and may reflect the presence of hydrocarbon sources remaining at the fire training pit. Elsewhere, the TOC concentrations did not exceed 25.5 mg/L. Six of seven TOC concentrations above 5 mg/L were located within the contaminant plume (Figure 4.10).

TOC concentrations measured at wells that are upgradient or crossgradient from the contaminant plume (e.g., monitoring wells TW101, MW107, and MW-116) averaged approximately 1.7 mg/L and are considered representative of background. The average background concentration of dissolved TOC is noteworthy because it represents additional organic matter that is available for use as a substrate in biodegradation reactions. The background concentrations likely represent carbon dissolved from organic matter dispersed throughout the aquifer. In addition to soil TOC, this carbon source should provide a continuing source of electron donors to fuel microbial redox reactions.

#### 4.4.1.2 Inorganic Chemistry

An interpretation of inorganic groundwater geochemistry, including isopleth maps for typical electron acceptors, can provide information regarding the relative importance of various biodegradation mechanisms and whether the conditions at the site are favorable for reductive dehalogenation of chlorinated compounds. Numerous

#### TABLE 4.7 GROUNDWATER GEOCHEMICAL PARAMETER ANALYTICAL DATA

							Total	
G 1 -	Dete	<b></b>			Total	Redox	Organic	Dissolved
Sample	Date	Temp		Conductivity	Alkalinity	Potential	Carbon	Hydrogen
ID	Sampled	(°C) <sup>a/</sup>	pН	(μS/cm) <sup>b/</sup>	(mg/L CaCO <sub>3</sub> ) <sup>c/</sup>	$(mV)^{d/}$	(mg/L) <sup>e/</sup>	(nM) <sup>f/</sup>
MW-105	5/15/97	18.9	5.3	77	20	79	22.2	0.35
MW-106	5/14/97	18.1	5.8	31	20	-100	1.93	0.14
MW-107	5/14/97	17.4	5.4	26	20	99	1.66	$NA^{g/}$
MW-111	5/15/97	19.0	4.8	19	10	222	1.37	5.79
MW-112	5/16/97	18.3	5.4	108	60	-184	22.0	3.67
MW-112A	5/16/97	18.8	5.4	19	10	200	0.884	19.02
MW-113	5/16/97	16.4	4.5	17	10	165	1.43	NA
MW-114	5/15/97	16.9	4.9	9	20	71	2.78	NA
MW-115	5/15/97	17.3	6.0	211	70	-201	8.59	0.1
MW-115A	5/15/97	19.1	4.8	24	10	196	1.06	0.3
MW-116	5/14/97	21.5	4.6	50	20	242	1.05	0.38
MW-116A	5/14/97	20.0	5.0	190	10	268	0.909	6.26
MW-117	5/15/97	19.9	4.4	62	10	319	3.72	0.1
MW-118	5/15/97	17.5	5.8	78	30	174	5.71	0.04
TW-101	5/14/97	21.8	6.2	124	50	74	2.39	1.13
TW-120	5/17/97	16.6	4.9	57	NA	60	NA	NA
TW-121	5/17/97	17.1	6.1	353	130	-65	25.5	NA
MPA	5/16/97	19.6	6.1	589	250	-56	144	NA
MPB	5/16/97	19.4	6.0	79	30	-35	5.27	NA
MPC	5/16/97	17.4	5.0	35	30	58	7.75	NA
MPD	5/16/97	18.1	5.6	43	NA	25	NA	NA
MPE	5/17/97	16.6	6.4	58	30	-7	1.51	NA
TMP-1	5/17/97	17.1	4.3	19	10	260	1.74	NA
TMP-2	5/17/97	17.8	5.1	57	20	-20	2.85	NA
TMP-3	5/17/97	16.5	4.6	40	10	255	1.76	NA
TMP-4	5/16/97	17.2	4.3	72	20	-5	2.71	NA
TMP-5	5/16/97	16.2	4.4	47	30	-30	NA	NA

a/ °C = Degrees Celsius.

b/ mS/cm = Microsiemens per centimeter.

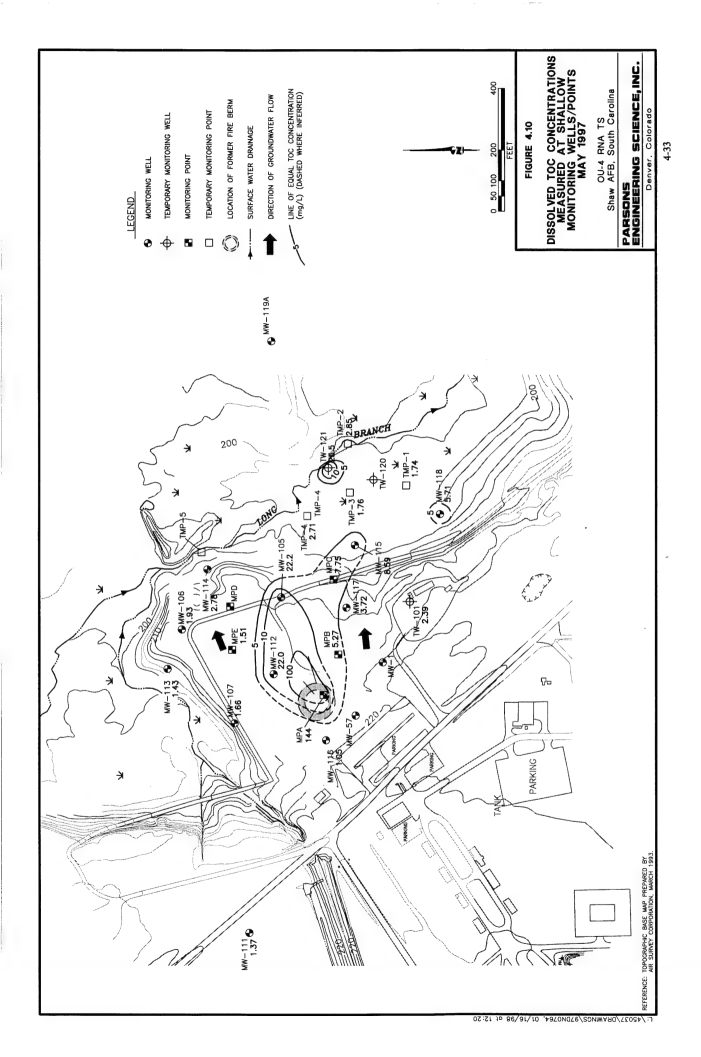
c/ CaCO<sub>3</sub> = Calcium Carbonate.

 $<sup>^{</sup>d'}$  mV = Millivolts.

e/ mg/L = Milligrams per liter.

<sup>&</sup>lt;sup>f/</sup> nM = Nanomoles.

g/ NA = Not available.



geochemical parameters were evaluated in site groundwater samples. Analytical results for geochemical parameters measured in site samples are presented in Table 4.6. The following paragraphs discuss those parameters that are most useful in the evaluation of site biodegradation processes.

#### 4.4.1.2.1 Dissolved Oxygen

DO concentrations were measured at monitoring wells and points during the May 1997 sampling event. Concentrations ranged from <0.1 to 9.0 mg/L (Table 4.6), with the highest concentrations generally measured in wells/points outside of the area of the contaminant plumes (Figure 4.11). The maximum DO concentration was measured at MW-117, which is unusual considering the presence of BTEX compounds at this location. It is possible that the chemical oxidation pilot testing performed at well MW-117 in November 1996 (IT, 1997) continued to affect groundwater chemistry in this area. At least 2,000 gallons of hydrogen peroxide was injected into monitoring well MW-117, along with quantities of ferrous sulfate and sulfuric acid to promote rapid oxidation of chlorinated compounds. Hydrogen peroxide was observed to immediately degrade upon introduction to form 0.5 mole of oxygen and 1 mole of water per mole of hydrogen peroxide degraded (IT, 1997). Therefore, the elevated DO concentration at MW-117 may be a residual effect of the pilot test.

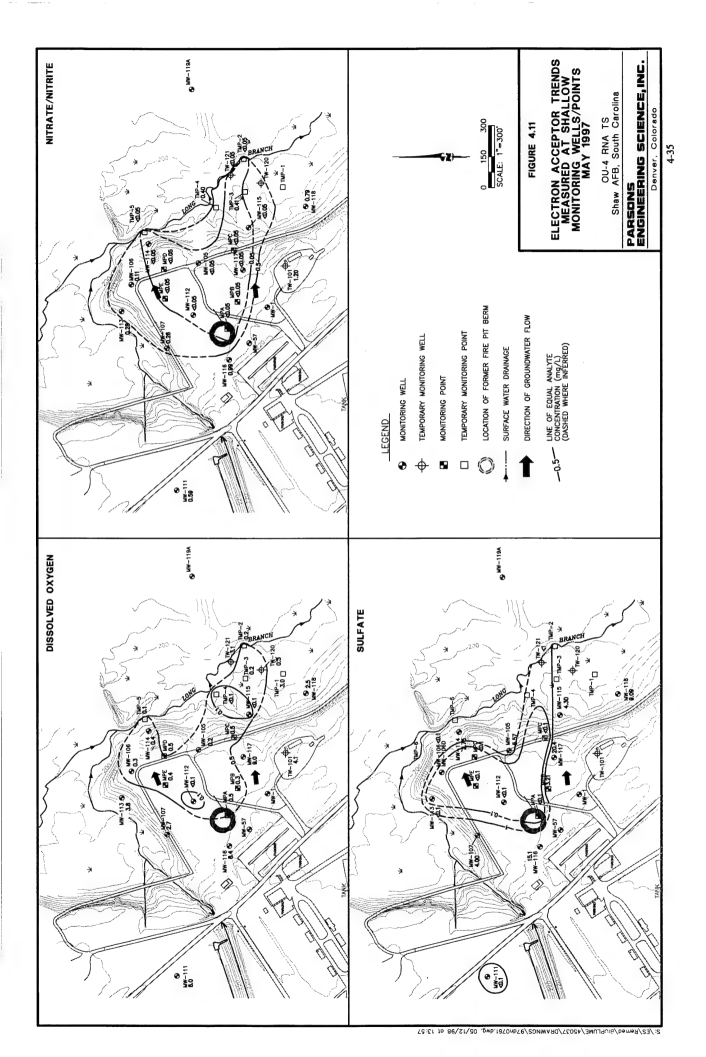
Throughout both plume lobes, most DO concentrations were generally less than 0.5 mg/L. The low concentrations of oxygen in the plume area relative to upgradient areas is a strong indication of anaerobic biological activity in the plume area. Depletion of DO within the BTEX and chlorinated solvent plumes at Site OU-4 indicates that microbial activity stimulated by the presence of anthropogenic hydrocarbons (e.g., BTEX, chlorobenzenes, VC, DCA, and DCE) and/or natural organic carbon is depleting DO and creating anaerobic conditions.

#### 4.4.1.2.2 Nitrate/Nitrite

Concentrations of nitrate/nitrite as nitrogen (N) were measured in groundwater samples collected in May 1997. Table 4.6 summarizes measured nitrate/nitrite (as N) concentrations, and Figure 4.11 shows the distribution of nitrate/nitrite (as N) concentrations in site groundwater. Background concentrations immediately upgradient from the plume ranged from about 0.8 mg/L to 1.2 mg/L. Within the plume, nitrate concentrations were generally less than 0.05 mg/L, whereas background concentrations averaged approximately 1.0 mg/L. The area of reduced nitrate concentrations also coincides with the area of reduced DO concentrations, indicating that as DO is removed, microbes are utilizing nitrate as an electron acceptor for degradation of organic material (such as BTEX or native organic carbon).

#### 4.4.1.2.3 Sulfate

Sulfate concentrations were measured in groundwater samples collected in May 1997. Sulfate concentrations in shallow groundwater at the site ranged from <0.1 mg/L to 20.4 mg/L (Table 4.6). Although the sulfate concentration at background well MW-116 exceeded 15 mg/L, sulfate exhibited high variability at the site, such that any sulfate concentrations exceeding 1 mg/L could potentially be representative of background. At the core of both lobes of the contaminant plume,



nearly all sulfate concentrations were less than 0.1 mg/L (Figure 4.11). The high sulfate concentration detected at MW-117 (20.4 mg/L) is believed to be a residual effect of the addition of sulfuric acid and ferrous sulfate during the chemical oxidation pilot test conducted at this location (IT, 1997).

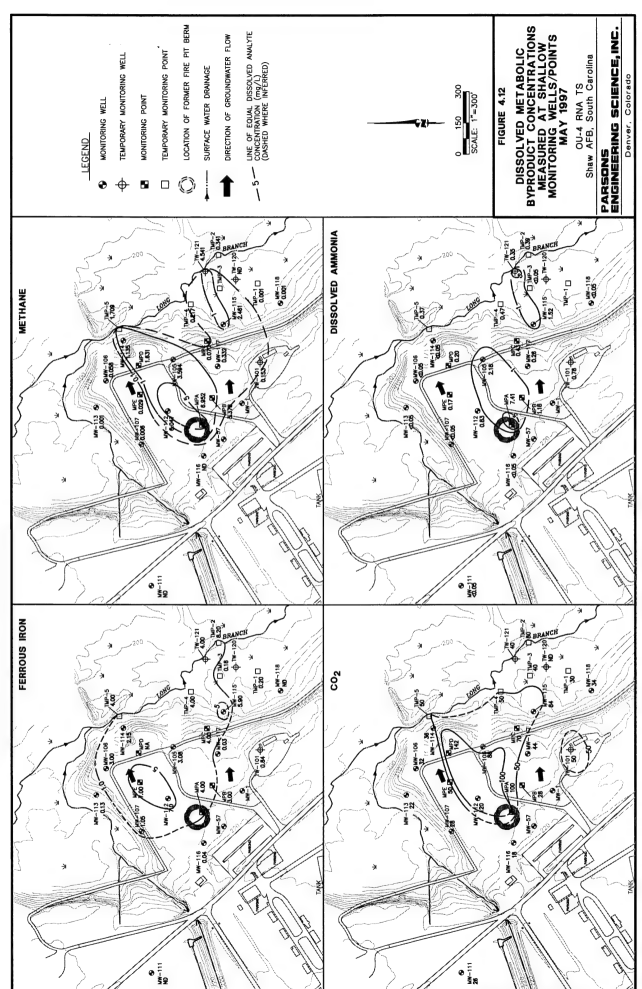
The area of decreased sulfate concentrations correlates well with chlorinated solvent and BTEX concentrations. It is therefore likely that sulfate reduction is an ongoing anaerobic biodegradation process at OU-4. This is important because when a system is sufficiently reducing for sulfate reduction to occur, conditions become more favorable for reductive halogenation of chlorinated solvents.

#### 4.4.1.2.4 Ferrous Iron

Ferrous iron [iron(II)] concentrations were measured in groundwater samples collected in May 1997. Table 4.6 summarizes ferrous iron concentrations. Measured ferrous iron concentrations range from 0.01 mg/L to 8.2 mg/L. Figure 4.12 shows the areal extent of ferrous iron in groundwater. Elevated ferrous iron concentrations coincide with both lobes of the solvent and BTEX plumes, suggesting that ferric iron hydroxide [iron(III)] is being reduced to ferrous iron during biodegradation of native organic carbon, BTEX compounds, and possibly less-chlorinated solvents. Background levels of ferrous iron average approximately 0.44 mg/L, as measured at upgradient and crossgradient wells (e.g., MW-116 and TW-101). The maximum ferrous iron concentrations (>5.0 mg/L) were detected in the vicinity of, or downgradient from, the zone of highest solvent or BTEX contamination in each of the two contaminant plume lobes (compare Figures 4.4 through 4.8 and 4.12).

Ferrous iron was not detected at elevated concentrations at MW-117 where significant chlorinated solvent and hydrocarbon contamination is present, and where large quantities of ferrous iron were added to groundwater during a chemical oxidation pilot test conducted 6 months prior to natural attenuation sampling (IT, 1997). Approximately 1,650 gallons of ferrous sulfate solution were added to the groundwater to help catalyze the oxidation of dissolved contaminants. Much of the ferrous iron likely was converted to ferric iron oxyhydroxides (Fe<sup>3+</sup>) when combined with oxygen released through hydrogen peroxide degradation. Ferrous iron is known to react instantly with oxygen in the dissolved phase to form ferric iron oxyhydroxide (Chappelle, 1993).

Recent evidence suggests that the reduction of ferric iron to ferrous iron cannot proceed at all without microbial mediation (Lovley and Phillips, 1988; Lovley et al., 1991; Chapelle, 1993). None of the common organic compounds found in low-temperature (i.e., non-hydrothermal), neutral, reducing groundwater could reduce ferric oxyhydroxides to ferrous iron under sterile laboratory conditions (Lovley et al., 1991). This means that the reduction of ferric iron requires microbial mediation by microorganisms with the appropriate enzymatic capabilities. Because the reduction of ferric iron cannot proceed without microbial intervention, the elevated concentrations of ferrous iron that were measured in the contaminated groundwater at the site are very strong indicators of microbial activity.



#### 4.4.1.2.5 Methane

Methane concentrations were measured in groundwater samples collected in May 1997. Table 4.6 lists methane concentrations, which ranged from 0.001 mg/L to 6.95 mg/L at the site. Figure 4.12 shows the distribution of methane in shallow site groundwater. The area of elevated methane concentrations correlates well with the chlorinated solvent and BTEX plumes. The presence of methane in the plume area indicates that conditions are sufficiently reducing for petroleum hydrocarbons and native organic matter to be used to support methanogenesis. Furthermore, the fact that methanogenesis is ongoing indicates that conditions in the plume area are strongly reducing, and therefore are favorable for reductive dehalogenation of chlorinated solvents.

Because methanogenesis produces more carbon dioxide than it consumes, an unlimited supply of carbon dioxide is theoretically available once the process of methanogenesis has been initiated. Therefore, methanogenesis is limited by the rate of reaction rather than the source of electron acceptors.

#### 4.4.1.2.6 Ammonia/Ammonium

The presence of ammonia/ammonium in groundwater can result from either nitrate reduction (facilitated by microbes) or fixing of atmospheric nitrogen (also a microbial process). Therefore, the presence of ammonia/ammonium in groundwater is a strong indication of microbial activity. Figure 4.12 shows the distribution of ammonia and ammonium (as N) in groundwater at OU-4. Ammonia/ammonium concentrations ranged from below detection limits (<0.05 mg/L) to 7.41 mg/L (Table 4.6), with the area of highest concentrations coinciding with the petroleum hydrocarbon source area at the fire training pit (MPA). A second area of high concentration was detected at TW-121, and is possibly the result of natural organic matter degradation near the ditch. Notably, ammonia/ammonium is present above 1 mg/L throughout the upgradient and downgradient portions of the east/southeast lobe of the contaminant plume, yet is present at lower concentrations through the central section of the plume. This may be an effect of the chemical oxidation pilot test performed at MW-117.

Given that the highest nitrate/nitrate (as N) concentration observed in site groundwater was 1.2 mg/L and that the highest ammonia/ammonium (as N) concentration is 7.41 mg/L, the nitrogen component of the ammonia process cannot entirely be derived from nitrate and may derive from atmospheric nitrogen gas. This implies that groundwater conditions within the BTEX and solvent plumes are reducing enough to support nitrogen fixation. Stumm and Morgan (1981) note that nitrogen fixation will occur when conditions are at sufficiently reducing to support iron reduction. This process can also occur when sulfate reduction or methanogenesis is favored. Therefore, this evidence of nitrogen fixation further confirms that conditions within the solvent and BTEX plumes are sufficiently reducing to allow reductive dehalogenation to proceed. The presence of ammonium further confirms that organic matter (anthropogenic or natural) is being biodegraded in the plume area.

#### 4.4.1.3 Ethane/Ethene in Groundwater

Ethene and ethane are the ultimate end products of the reductive dehalogenation of PCE, TCE, and 1,1,1-TCA. Concentrations of ethane and/or ethene below 0.01 mg/L were detected along the east/southeast contaminant plume lobe and at TMP-5 in the toe of the northeast contaminant plume. Figure 4.13 shows ethene and ethane distributions, and Table 4.6 lists detected concentrations. The largest concentrations of ethane and ethene emanated from the suspected source area at MPC. Although ethene and ethane occurred at very low concentrations, the presence of these compounds at and downgradient from an identified solvent source area is a strong indicator of reductive dehalogenation.

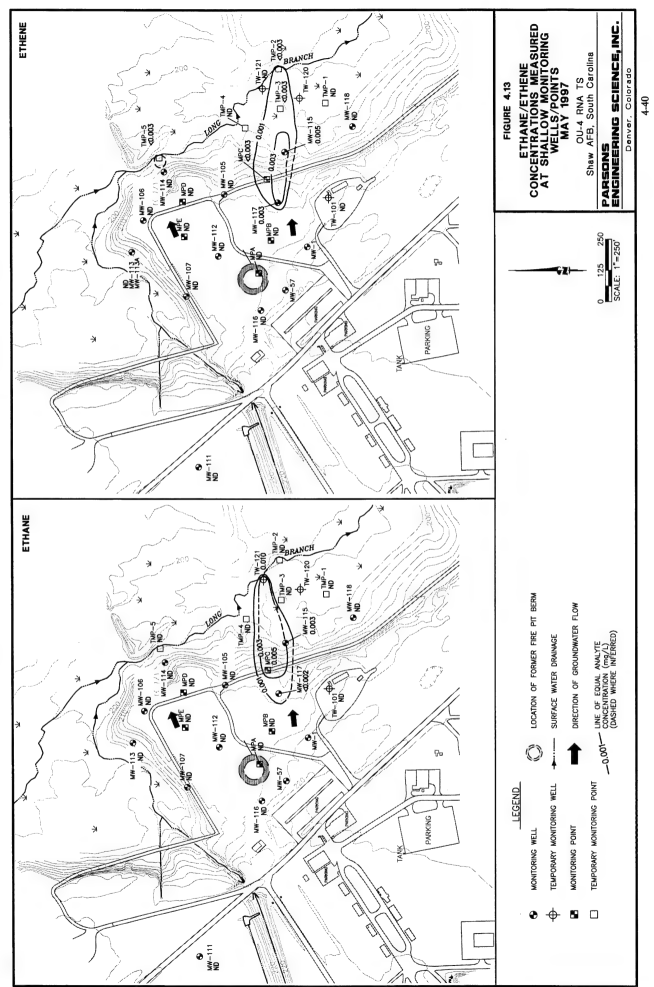
#### 4.4.2 Additional Geochemical Indicators

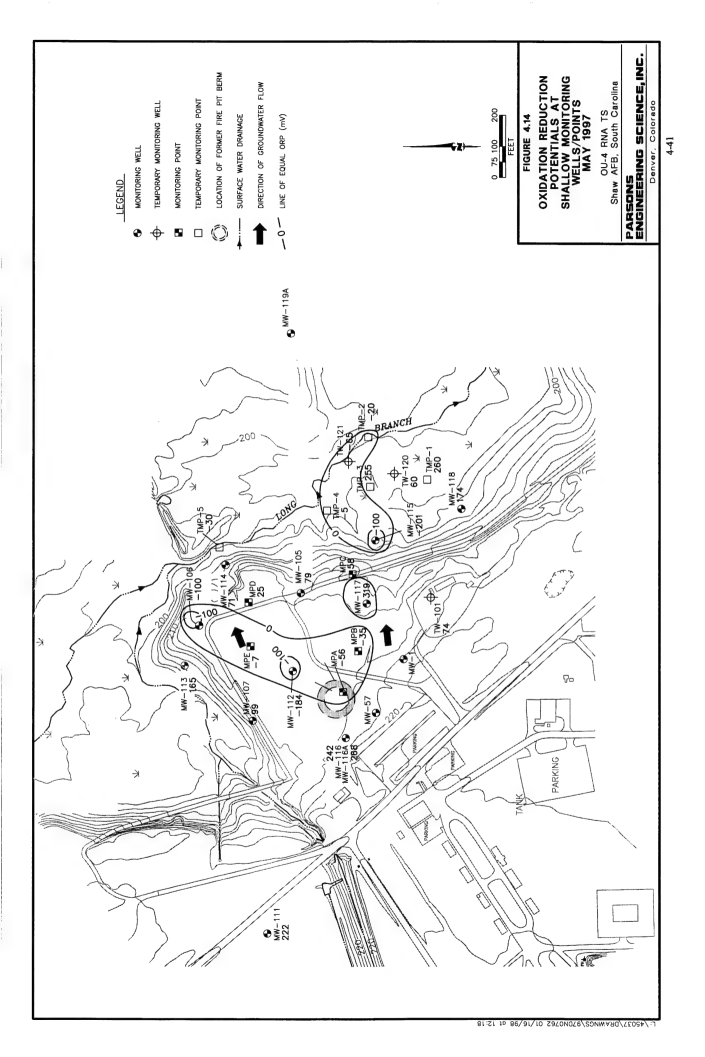
Other geochemical data collected for this evaluation can be used to further interpret and support the contaminant, electron donor, electron acceptor, and byproduct data previously discussed. These parameters provide additional qualitative indications of what biodegradation processes may be operating at the site.

### 4.4.2.1 Oxidation/Reduction Potential and Dissolved Hydrogen as Indicators of Redox Processes

ORPs were measured at groundwater monitoring wells and points in May 1997. ORP is a measure of the relative tendency of a solution to accept or transfer electrons. The ORP of a groundwater system depends on which electron acceptors are being reduced by microbes during oxidation of organic compounds. ORPs for shallow groundwater at the site ranged from 319 millivolts (mV) to -201 mV. summarizes available ORP data, and Figure 4.14 illustrate ORP trends. Low ORPs (<0 mV) coincide well with the northeast contaminant plume; ORPs below 0 are present through the upgradient and downgradient portions of the east/southeast contaminant plume. The higher ORPs through the central portion of the east/southeast contaminant plume (MPC and MW-117) may have resulted from the chemical oxidation pilot test, which caused a large release of DO (IT, 1997). Elsewhere, however, the correlation between ORP and contaminant plume extent; decreased DO, nitrate, and sulfate concentrations; and elevated ferrous iron and methane concentration is For example, MW-106 had a relatively low ORP (-100 mV), but inconsistent. relatively low BTEX and solvent concentrations were detected at this location. Unexplained inconsistencies may result from mis-measurement of ORP due to sampling error, varying instrument electrode sensitivities to different redox couples, aquifer heterogeneity, or screening over multiple aquifer subunits.

Concentrations of dissolved hydrogen (H<sub>2</sub>) also can be used to evaluate redox processes in groundwater systems (Lovley and Goodwin, 1988; Lovley et al., 1994; Chapelle et al., 1995). H<sub>2</sub> is continuously produced in anaerobic groundwater systems by fermentative microorganisms that decompose natural and anthropogenic organic matter. This H<sub>2</sub> is then consumed by respiratory microorganisms that use nitrate, ferric iron, sulfate, or CO<sub>2</sub> as terminal electron acceptors. These nitrate-, ferric iron-, sulfate- and CO<sub>2</sub>-reducing (methanogenic) microorganisms exhibit different efficiencies in utilizing the H<sub>2</sub> that is being continually produced. Nitrate reducers are highly efficient H<sub>2</sub> utilizers and maintain very low steady-state H<sub>2</sub> concentrations. Ferric iron reducers





are slightly less efficient and thus maintain somewhat higher H<sub>2</sub> concentrations. Sulfate reducers and methanogenic bacteria are progressively less efficient and maintain even higher H<sub>2</sub> concentrations. Because each terminal electron accepting process has a characteristic H<sub>2</sub> concentration associated with it, H<sub>2</sub> concentrations can be an indicator of predominant redox processes. These characteristic ranges are given in Table 4.8.

Dissolved H<sub>2</sub> concentrations measured in OU-4 groundwater in May 1997 are summarized in Table 4.7. Concentrations ranged from 0.1 nanomoles per liter (nM/L) to 5.79 nM/L at shallow monitoring wells. The two highest H2 concentrations were measured either far upgradient of the main CAH plume at well MW-111 (5.79 nM/L) or along the northeast contaminant plume lobe at MW-112 (3.67 nM/L). remaining H<sub>2</sub> values, which ranged from 0.1 to 1.13 nM/L, were obtained at wells scattered across the site, with no apparent correlation to groundwater contamination. Most elevated dissolved H2 concentrations were not located in zones of high groundwater contamination as would be expected. The magnitude of dissolved hydrogen values at locations within the plume suggest that denitrification, ferric iron reduction, and sulfate reduction should be possible at shallow monitoring wells/point intervals in the plume. Dissolved H2 concentrations measured in excess of 5 nM/L at the site (though not in the groundwater plume) indicate that methanogenic conditions can be achieved at the site. A dissolved hydrogen concentration of 19.02 nM/L was measured at MW-112A which is suggestive of highly reducing methanogenic conditions. No contamination was detected in monitoring well MW-112A.

TABLE 4.8

RANGE OF HYDROGEN CONCENTRATIONS FOR A GIVEN TERMINAL ELECTRON-ACCEPTING PROCESS OU-4 RNA TS
SHAW AFB. SOUTH CAROLINA

Terminal Electron- Accepting Process	Dissolved Hydrogen Concentration (nanomoles per liter)
Denitrification	< 0.1
Ferric Iron Reduction	0.2 to 0.8
Sulfate Reduction	1 to 4
Methanogenesis	5 to 20

#### 4.4.2.2 Alkalinity and Carbon Dioxide Evolution

Carbon dioxide is produced during the biodegradation of anthropogenic and native organic carbon compounds. Carbon dioxide forms carbonic acid that dissolves carbonate minerals where present in the aquifer matrix, thereby increasing the alkalinity of the groundwater. An increase in alkalinity [measured as calcium carbonate (CaCO<sub>3</sub>)] in an area with BTEX or chlorinated solvent (particularly less-chlorinated solvents)

concentrations elevated above background conditions can be used to infer that petroleum hydrocarbons, less-chlorinated solvents, or native organic carbon have been consumed through aerobic and anaerobic microbial respiration.

Total alkalinity (as CaCO<sub>3</sub>) was measured in groundwater samples collected in May 1997. These measurements are summarized in Table 4.7. Alkalinity is a measure of the ability of groundwater to buffer changes in pH caused by the addition of biologically generated acids. Total alkalinity at the site varied from 10 mg/L to 250 mg/L, with many of the highest concentrations measured in wells/points corresponding to the presence of significant solvent or BTEX contamination (e.g., monitoring wells/points MW-112, MW-115, and MPA). However, alkalinity within the plume area is variable. The limited mass of carbonates is supported by a low natural groundwater alkalinity (often as low as 10 mg/L), elevated carbon dioxide concentrations within the plume as discussed below, and low groundwater pH as described in Section 4.4.2.4.

Free carbon dioxide concentrations also were measured in groundwater samples collected in May 1997 (Table 4.6). As shown on Figure 4.12, carbon dioxide concentrations within the full extent of the chlorinated solvent and BTEX plumes are noticeably elevated above background concentrations. Background concentrations are about 18 to 50 mg/L, while concentrations within the plume range from 50 to 120 mg/L. Carbon dioxide concentrations above background result from both aerobic and anaerobic biodegradation processes as contaminants are ultimately converted to carbon dioxide and water.

#### 4.4.2.4 pH

pH was measured for groundwater samples collected from monitoring points and monitoring wells in May 1997 (Table 4.7). The pH of a solution is the negative logarithm of the hydrogen ion concentration [H<sup>+</sup>]. The groundwater pH measured at the site ranged from 4.3 to 6.4 standard units, which is slightly below the optimal range for most microbial populations that degrade organic matter. The slightly acidic pH measurements indicates that microbial reactions and the resultant production of organic acids by microbial degradation of hydrocarbons may have an effect on groundwater pH. The moderate to low alkalinity of site groundwater is consistent with low observed pH readings, and suggests a limited capacity of groundwater to buffer the generation or organic acids. However, low pHs also were observed in upgradient wells, suggesting generally low pH readings across the entire aquifer.

#### 4.4.2.5 Temperature

Groundwater temperature was measured at groundwater monitoring points and monitoring wells in May 1997. Table 4.7 summarizes groundwater temperature readings. Temperature can affect the types and growth rates of bacteria that can be supported in the groundwater environment, with higher temperatures typically resulting in higher growth rates. Groundwater temperatures at OU-4 varied from 16.2 degrees Celsius (°C) to 21.8°C. These relatively warm temperatures indicate that microbially-mediated biodegradation reactions should not be inhibited by overly cold groundwater temperatures, and may be enhanced.

#### 4.5 APPROXIMATION OF BIODEGRADATION RATES

Biodegradation rate constants are necessary to accurately simulate the fate and transport of contaminants dissolved in groundwater. In many cases, biodegradation of contaminants can be approximated using first-order kinetics. First-order biodegradation rate constants may be calculated on the basis of field-scale data, or by using a small sample of the aquifer material and groundwater for microcosm studies.

For reductive dehalogenation of chlorinated compounds, first-order rates are not necessarily an accurate representation because the degradation of CAHs depends on both the concentrations of electron donors (e.g., fuel hydrocarbons or native organic material) and the concentrations of electron acceptors (including inorganic acceptors and CAHs) (Moutoux et al., 1996). Second-order rates would therefore allow more accurate modeling, but at this time there are no readily available groundwater contaminant transport models that can utilize such rates. Therefore, first-order rates must be estimated for such an application.

Microcosm studies are used to show that the microorganisms necessary for biodegradation are present and to help quantify rates of biodegradation. Because microcosm studies are time consuming and expensive, they should be undertaken only at sites where there is considerable skepticism concerning the biodegradation of contaminants. The results of a microcosm study are strongly influenced by the nature of the geological material submitted for study, the physical properties of the microcosm, the sampling strategy, and the duration of the study. Biodegradation rate constants determined by microcosm studies often are much higher than those observed in the field. Microcosms are most appropriate as indicators of the potential for natural bioremediation, and to prove that losses are biological, but it may be inappropriate to use them to generate rate constants. The preferable method of contaminant biodegradation rate-constant determination is by use of field data.

In order to calculate first-order biodegradation rate constants on the field scale, the apparent degradation rate must be normalized for the effects of dilution, dispersion, and sorption. One method proposed by Buscheck and Alcantar (1995) involves interpretation of a steady-state contaminant plume and is based on the one-dimensional, steady-state analytical solution to the advection-dispersion equation presented by Bear (1979). This method can be adapted to estimate rate constants for CAHs. Another method for estimating dehalogenation rates for CAHs is described by Moutoux et al. (1996). This method provides a total dechlorination rate for all dechlorination steps. All rates (including the rapid TCE to DCE rate and the slow VC to ethene) are averaged together using the Moutoux et al. (1996) method.

The Buscheck and Alcantar (1995) and Moutoux *et al.* (1996) methods were used to estimate first-order biodegradation rate constants for BTEX and chlorinated solvents at OU-4. The methods and related calculations are summarized in Table 4.9. Decay rates

SUMMARY OF CHLORINATED SOLVENT AND BTEX DECAY RATES SHAW AFB, SOUTH CAROLINA **OU-4 RNA TS** TABLE 4.9 MAY 1997

Analysis Method	Contaminants	Dechlorination Sequence	Sampling Date	Decay Rate (day <sup>-1</sup> )	Half Life (days)
B&A²∕	Chlorinated Ethene Plume	PCE - TCE - DCE (all isomers) - VC	May 1997	1.24 x 10 <sup>-2</sup>	99
Reductive Dechlorination <sup>b/</sup>	Chlorinated Ethene Plume	PCE - TCE - DCE (all isomers) - VC	May 1997	2.61 x 10 <sup>-3</sup>	566
B&A	Chlorinated Ethane Plume	1,1,1-TCA - 1,1-DCA	May 1997	$1.31 \times 10^{-2}$	53
Reductive Dechlorination	Chlorinated Ethane Plume	1,1,1-TCA - 1,1-DCA	May 1997	$2.49 \times 10^{-3}$	213
B&A	Chlorinated Benzene Plume	1,2-DCB + 1,3-DCB + 1,4-DCB - CB	May 1997	$1.13 \times 10^{-2}$	61
Reductive Dechlorination	Chlorinated Benzene Plume	1,2-DCB + 1,3-DCB + 1,4-DCB - CB	May 1997	NA	NA
B&A	BTEX Plume	NA	May 1997	$1.40 \times 10^{-2}$	20
Reductive Dechlorination	BTEX Plume	NA	May 1997	NA	NA

B&A = Method of Buscheck and Alcantar (1995) for steady-state plumes.

4-45

b/ Reductive Dechlorination = Method of Moutoux et al. (1996), which gives decay rate attributable to reductive dechlorination. Note: The flow path chosen to calculate the biodegradation rates is MPC to MW-115 to TMP-3. The average total organic carbon content of soils was assumed to be 0.00024 mg/mg (Rust E&I, 1995) for retardation rates used in the calculation of biodegradation rates (see Appendix D). computed using data from Site OU-4 and the Buscheck and Alcantar (1995) method were 0.014 day<sup>-1</sup> for total BTEX, 0.012 day<sup>-1</sup> for total chlorinated ethenes, 0.013 day<sup>-1</sup> for total chlorinated ethane (1,1,1-TCA to DCA isomers only), 0.011 day<sup>-1</sup> for dichlorobenzenes to CB. The equivalent half-lives are 50 days, 56 days, 53 days, and 61 days, respectively. Decay rates computed using the Moutoux *et al.* (1996) method indicated reductive dechlorination rates of 0.0026 day<sup>-1</sup> (half-life of 266 days) for total chlorinated ethene contamination and 0.0025 day<sup>-1</sup> (half-life of 213 days) for chlorinated ethane contamination.

#### 4.6 DISCUSSION

Compounds detected in groundwater at OU-4 include BTEX, PCE, TCE, cis-1,2-DCE, trans-1,2-DCE, 1,1-DCE, VC, 1,1,1-TCA, 1,1-DCA, chloroform, 1,2-DCB, 1,3-DCB, 1,4-DCB, and CB. Of these compounds, benzene, toluene, PCE, TCE, cis-1,1-DCE, 1,1-DCE, VC, and 1,1,1-TCA were detected at concentrations exceeding USEPA MCLs. 1,1-DCA was detected at a maximum concentration of 8,700 µg/L, but no federal MCL is available for this compound.

The most recent comprehensive groundwater sampling event prior to groundwater sampling performed for this TS occurred between March and May 1993. Contaminant concentrations in single monitoring wells generally have fluctuated over time, with no clear trends of increasing or decreasing concentrations within the primary source area (i.e., the fire training pit). This is likely a function of the low concentrations, mixing in wells during sampling, slight differences in sampling procedures, variations in plume concentrations due to water level/flow direction changes, and/or variable rates of desorption from aquifer solids.

The additional data collected for this effort helped to define several possible additional sources of groundwater contamination in the OU-4 area. In addition to the former fire training pit, these sources include the areas around monitoring points MPB and MPC. A slight product sheen was detected at MPD, but contaminant concentration isopleths do not suggest that a significant contaminant source exists at this location. A contaminant source was identified near MW-107 during the RI; however, no groundwater contamination was detected near this location during TS sampling, and previously detected soil contamination at this location may no longer be a threat. The presence of elevated soil and groundwater contaminant concentrations between the fire training pit and MPC suggests that contamination is spread as residual and/or mobile LNAPL across a narrow 500-foot corridor between these two locations. This soil contamination likely is the result of LNAPL migration on top of the water table, smearing with water level fluctuations, or the historical use of multiple fire training locations.

Chemical and geochemical evidence indicates that the contaminants at OU-4 are being biodegraded, either as substrates or as electron acceptors. Geochemical data indicate that DO, nitrate, and sulfate are being consumed, and iron (II), methane, and ammonia/ammonium are being produced within the dissolved plume. This evidence suggests that microbial consumption of anthropogenic and natural organic matter (carbon compounds) is taking place at OU-4. Anthropogenic carbon compounds are present as petroleum hydrocarbons and less-chlorinated solvents (including, but not limited to, BTEX, CB, VC, DCE, and DCA) that are dissolved in groundwater.

Native carbon compounds include solid organic matter in the aquifer matrix and whatever portion of that matter that has dissolved into site groundwater. These compounds can serve as electron donors in redox reactions that also are consuming electron acceptors (e.g., DO, ferric iron, carbon dioxide, and chlorinated hydrocarbons) and creating byproducts [e.g., methane, iron (II), and carbon dioxide]. Plots of contaminants, electron donors, electron acceptors, and byproducts of biodegradation reactions (Figures 4.4 through 4.13) provide strong evidence of these processes.

Microbial consumption of both native and anthropogenic organic carbon compounds destroys those compounds and creates conditions favorable for reductive dehalogenation. Site chemical data provide evidence of dehalogenation of TCE, DCE, TCA, and DCA. The presence of DCE and VC in site groundwater provides the primary line of chemical evidence for reductive dehalogenation of chlorinated ethenes. In addition, the relative abundance of the cis-1,2 isomer of DCE relative to the trans-1,2 and 1,1- isomers of DCE further supports chlorinated ethene reductive dehalogenation. The presence of 1,1-DCA also suggests that 1,1,1-TCA is being reductively dehalogenated in the east/southeast plume lobe. The presence of low concentrations of ethane and ethene further indicate that conditions are sufficiently reducing for the dehalogenation processes to proceed to completion.

Elevated chloride concentrations within the solvent plumes also may indicate dehalogenation reactions, although chloride can also be produced from oxidation of less-chlorinated CAHs (i.e., utilization as a substrate). Nonetheless, elevated chloride concentrations throughout the contaminant plume support the occurrence of CAH biodegradation.

Evidence also suggests that 1,1,1-TCA is abiotically dehydrohalogenated to 1,1-DCE in the east/southeast contaminant plume lobe. It appears that 1,1-DCE concentrations coincide with significant 1,1,1-TCA concentrations, but that TCE concentrations are relatively low at these same locations. On the other hand, the presence of 1,1-DCA in wells that also contained BTEX and chlorinated ethenes suggests that reductive dechlorination of TCE and TCA is ongoing within the east/southeast contaminant plume lobe. It therefore appears that 1,1,1-TCA may be both reductively dehalogenated to 1,1-DCA and abiotically transformed to 1,1-DCE.

The ocurrence of CB and DCB contamination indicates that some reductive dechlorination of DCB may be occurring. However, DCB and CB, which are the least chlorinated of the chlorobenzenes, are also the least susceptible to reductive dechlorination. Therefore, it is possible that both CB and DCB are source solvents, and that contaminant mass reductions may be occurring through aerobic reactions where the chlorobenzenes act as electron donors/substrates in a manner similar to BTEX biodegradation.

Wiedemeier et al. (1996b) present a worksheet to allow an initial assessment of the prominence of reductive dehalogenation at a site. The worksheet, including the point values determined for OU-4, is included as Table 4.10. The interpretation of points awarded during the screening process is shown in Table 4.11. The score for OU-4 computed using Table 4.10 is 27, indicating that strong evidence for reductive dehalogenation of chlorinated organics is present.

#### TABLE 4.10 ANALYTICAL PARAMETERS AND WEIGHTING FOR PRELIMINARY SCREENING

Analysis	Concentration in Most Contaminated Zone	Interpretation	Value	OU-4 Score
Oxygen	<0.5 mg/L	Tolerated, suppresses the reductive pathway at higher concentrations	3	3
Oxygen	>1 mg/L	VC may be oxidized aerobically	-3	
Nitrate	<1 mg/L	At higher concentrations may compete with reductive pathway	2	2
Iron II	>1 mg/L	Reductive pathway possible	3	3
Sulfate	<20 mg/L	At higher concentrations may compete with reductive pathway	2	2
Sulfide	>1 mg/L	Reductive pathway possible	3	0
Methane	<0.5 mg/L	VC oxidizes	0	3
	>0.5 mg/L	Ultimate reductive daughter product, VC Accumulates	3	
Oxidation	< 50 millivolts (mV)	Reductive pathway possible	1	1
Reduction Potential (redox potential)	<-100mV	Reductive pathway likely	2	
pН	5 < pH < 9	Optimal range for reductive pathway	0	0
	5 > pH >9	Outside optimal range for reductive pathway	-2	
TOC	> 20 mg/L	Carbon and energy source; drives dechlorination; can be natural or anthropogenic	2	0
Temperature	> 20°C	At T >20°C biochemical process is accelerated	I	0
Carbon Dioxide	>2x background	Ultimate oxidative daughter product	1	1
Alkalinity	>2x background	Results from interaction of carbon dioxide with aquifer minerals	1	0
Chloride	>2x background	Daughter product of organic chlorine	2	2
Hydrogen	>1 nM	Reductive pathway possible, VC may accumulate	3	0
Hydrogen	<1 nM	VC oxidized	0	0

# TABLE 4.10 (Concluded) ANALYTICAL PARAMETERS AND WEIGHTING FOR PRELIMINARY SCREENING

Analysis	Concentration in Most Contaminated Zone	Interpretation	Value	OU-4 Score
Volatile Fatty Acids	> 0.1 mg/L	Intermediates resulting from biodegradation of aromatic compounds; carbon and energy source	2	0
BTEX	> 0.1 mg/L	Carbon and energy source; drives dechlorination	2	2
PCE		Material released	0	0
TCE <sup>a/</sup>		Material released Daughter product of PCE	0 2 a/	0
DCE <sup>a</sup>		Material released Daughter product of TCE. If cis is greater than 80% of total DCE it is likely a daughter product of TCE	0 2 <sup>a/</sup>	0 2
VC <sup>a/</sup>		Material released Daughter product of DCE	0 2 a/	0 2
Ethene/Ethane	>0.01mg/L >0.1 mg/L	Daughter product of VC/ethene	2 a/ 3 a/	0
Chloroethane		Daughter product of VC under reducing conditions	2 a/	0
1,1,1- Trichloroethane		Material released	0	0
1,2- Dichlorobenzene		Material released	0	0
1,3- Dichlorobenzene		Material released	0	0
1,4- Dichlorobenzene		Material released	0	0
Chlorobenzene		Material released or daughter product of dichlorobenzene	2 a/	2
1,1-DCE		Daughter product of TCE or chemical reaction of 1,1,1-TCA	2 a/	2
			Total	27

a/ Points awarded only if it can be shown that the compound is a daughter product (i.e., not a constituent of the source NAPL).

#### TABLE 4.11 INTERPRETATION OF POINTS AWARDED DURING NATURAL ATTENUATION SCREENING

#### OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

Score	Interpretation
0 to 5	Inadequate evidence for biodegradation of chlorinated organics
6 to 14	Limited evidence for biodegradation of chlorinated organics
15 to 20	Adequate evidence for biodegradation of chlorinated organics
>20	Strong evidence for biodegradation of chlorinated organics

Given the available evidence, the dissolved CAH plume at OU-4 exhibits characteristics of both type 1 and type 3 behavior. Dissolved petroleum hydrocarbons and possibly some less-chlorinated solvents appear to act as carbon sources. It is likely that the introduction of petroleum hydrocarbons and solvents due to fire training activities stimulated microbial activity and made the groundwater system reducing enough to allow reductive dehalogenation of TCE, DCE, TCA, and VC. VC has been created due to dehalogenation, but it is possible that it is being oxidized (as a substrate) once it enters aerobic groundwater, especially near Long Branch Creek. Likewise, it is possible that DCE and DCA are oxidized as well. However, conclusive evidence of these oxidation processes has not been gathered. It is possible that some microbial consumption of native organic matter is taking place outside the plume and within the plume, but the DO values measured outside of the contaminant plume suggest this process is minor. Type 2 behavior may therefore be limited in extent and is not discernible at OU-4.

Maximum rates of BTEX, CB, chlorinated ethene, and chlorinated ethane degradation estimated from data collected for this investigation were 0.014 day<sup>-1</sup>, 0.012 day<sup>-1</sup>, 0.013 day<sup>-1</sup>, and 0.011 day<sup>-1</sup>, respectively. As petroleum hydrocarbons and native organic matter continue to be consumed, reductive dehalogenation of the chlorinated ethenes will continue; however, once BTEX compounds are completely degraded, the rate at which dehalogenation proceeds may slow as the microbes utilize less easily degradable fuel hydrocarbons or other organic matter.

As the type of organic matter used during biodegradation changes, the ORP conditions also may change and alter which reactions are favored. This could also affect the rates at which CAHs are reductively dehalogenated, and could also inhibit dehalogenation of some compounds. If and how this will happen is difficult to predict; the relative "degradability" of the available organic carbon and the effect of a changing electron donor source is not yet well understood. It can be stated that if conditions become more oxidizing (but not aerobic), the dehalogenation of DCE may also slow or cease. However, aerobic and/or anaerobic mineralization of VC and DCE may become more important (i.e., type 3 conditions may dominate).

#### **SECTION 5**

## GROUNDWATER FLOW AND CONTAMINANT TRANSPORT MODEL

#### 5.1 GENERAL OVERVIEW AND MODEL DESCRIPTION

In order to help predict the future migration and fate of CAHs dissolved in groundwater at OU-4, Parsons ES modeled shallow groundwater flow and the fate and transport of the dissolved CAH plume. The modeling effort had three primary objectives: 1) predict the future extent and concentrations of the dissolved contaminant plume by modeling the combined effects of advection, dispersion, sorption, and biodegradation; 2) assess the potential for downgradient receptors to be exposed to contaminants at concentrations above regulatory levels of concern; and 3) if applicable, provide technical support for the natural attenuation remedial option at post-modeling regulatory negotiations. The model was developed using site-specific data and conservative assumptions about governing physical and chemical processes. The model input data are believed to be reasonably conservative so that the actual reduction in contaminant mass caused by natural attenuation should meet or exceed model predictions.

The model codes MODFLOW (McDonald and Harbaugh, 1988) and MT3D<sup>96®</sup> (S.S. Papadopulos & Associates, Inc., 1996) were used to estimate the potential for dissolved CAH migration and degradation by naturally occurring mechanisms operating at the site. MODFLOW was used to generate a groundwater flow model for the site. The MODFLOW flow field was incorporated into the transport solution computed by  $MT3D^{96}$  (version 1.1). The pre- and post- processors contained in Visual MODFLOW, version 2.20 (Waterloo Hydrogeologic Software, 1996) were used to facilitate model development and analysis and presentation of the model results. The MT3D<sup>96®</sup> code incorporates advection, dispersion, sorption, and biodegradation to simulate contaminant plume migration and degradation. MT3D<sup>96®</sup> uses solution routines based on the Method of Characteristics (MOC) solute transport model [e.g., as developed by Konikow and Bredehoeft (1978) for the USGS two-dimensional (2-D) MOC model code]. The MOC model was modified by Zheng (1990) to allow threedimensional (3-D) solutions, and to allow use of a modified MOC that reduces numerical dispersion; the modified model was called MT3D. MT3D% is an improvement on MT3D that allows for spatial variation of solute transport parameters (e.g., retardation and first-order decay rate).

Ideally, a code for simulating degradation of CAHs would track daughter products as well as parent compounds and allow specification of varying decay rates for each compound. Battelle National Laboratories may have developed and released such a code under the name RT3D by modifying MT3D.

#### 5.2 CONCEPTUAL MODEL DESIGN AND ASSUMPTIONS

On the basis of the data presented in Section 3, the shallow saturated zone was conceptualized and modeled as a homogeneous, unconfined aquifer. Hydrogeologic profiles of the site (Figure 3.2) show that the major subsurface units of the aquifer are comprised of Duplin Formation terrace deposits characterized by mostly medium to very coarse sands that become finer with increasing depth. The unconfined aquifer is bounded at depth by the "100-foot clay" of the Black Creek Formation that acts as an aquitard between the shallow aquifer and the underlying Lower Black Creek Aquifer. The CAH plume is migrating primarily within the upper surface of the aquifer, as evidenced by the general lack of CAH detections at three intermediate wells at the site (MW-112A, -115A, and -116A). An observed increase in the fine and clay content of the soils begins at approximately 30 to 35 feet bgs and limits the vertical movement of contaminants between shallow and intermediate well locations.

Depth to groundwater across the site ranges from 0 to 19 feet bgs, with depth to groundwater decreasing nearer to Long Branch Creek. The configuration of groundwater elevation isopleths on Figure 3.5 indicates that groundwater in the OU-4 area primarily flows northeast and east from the fire-training burm, with discharge to Long Branch Creek east of the site and an adjoining tributary to the north of the site. The average horizontal hydraulic gradient across the study area is approximately 0.007 ft/ft; however, the groundwater gradient increases to approximately 0.11 ft/ft below the upland terrace near the creek. Vertical flow profiles indicate downward flow (0.050 to 0.077 ft/ft) at well clusters across the site.

Groundwater elevations in May 1997 (Figure 3.5) generally were lower than groundwater elevations observed during previous investigations (by approximately 2 feet); however, groundwater flow directions at the site were consistent. Therefore, it was assumed that the groundwater migration patterns interpreted from May 1997 water levels were representative of steady-state conditions. It is assumed that recharge from precipitation is substantial because annual precipitation at the Base is approximately 46 inches, the ground surface adjacent to fire training area source locations is sparsely vegetated, and sandy soils permit rapid infiltration to a relatively shallow water table (Section 3).

For the purposes of the model, it is assumed that mobile and residual LNAPL (cosolvenated CAHs and fuel hydrocarbons) contributes CAH contamination to groundwater. Although the last reported use of the fire training pit occurred in 1969, groundwater velocities are sufficiently high that a particle of contamination can travel from the fire training berm to Long Branch Creek within 2 years. Therefore, the groundwater model was calibrated with the assumption that existing LNAPL sources provide a continuous source of contaminants to groundwater. It is likely that the source will weather over time, and the mass of contaminants entering groundwater will decline.

The most important assumption made when using the MT3D<sup>96®</sup> code is that dispersion, sorption and biodegradation are major factors controlling contaminant fate and transport at the site. According to soil TOC data previously collected at the site (Rust E&I, 1995), concentrations of organic carbon within the site soil may provide limited contaminant sorption. Data also suggest that substantial biodegradation of

CAHs is occurring within the plume. Dispersivity, which is a characteristic of the porous medium, is a measure of the longitudinal and lateral spreading of the contaminant plume caused by local heterogeneities that cause deviations from the average linear migration velocity. The magnitude of dispersion is generally believed to be scale-dependent: the longer the plume flowpath, the greater the dispersion. Given the intercepted plume flowpath by Long Branch Creek and the rapid groundwater velocities, it is reasonable to assume that dispersion is not an important parameter influencing solute transport in the study area. Dispersion is estimated using literature values and accepted rules-of-thumb. Sorption (assumed to be a linear process) is simulated using a coefficient of retardation, and biodegradation is simulated using a first-order decay constant. Selection of values for these model input parameters is discussed in Section 5.3.3.

#### 5.3 INITIAL MODEL SETUP

Where possible, the initial setup for this model was based on site-specific data. Where site-specific data were not available (e.g., effective porosity), reasonable assumptions were made on the basis of widely-accepted literature values for materials similar to those found in the shallow aquifer at OU-4. The following sections describe the basic model setup. Those model parameters that were varied during model calibration are discussed in Section 5.4.

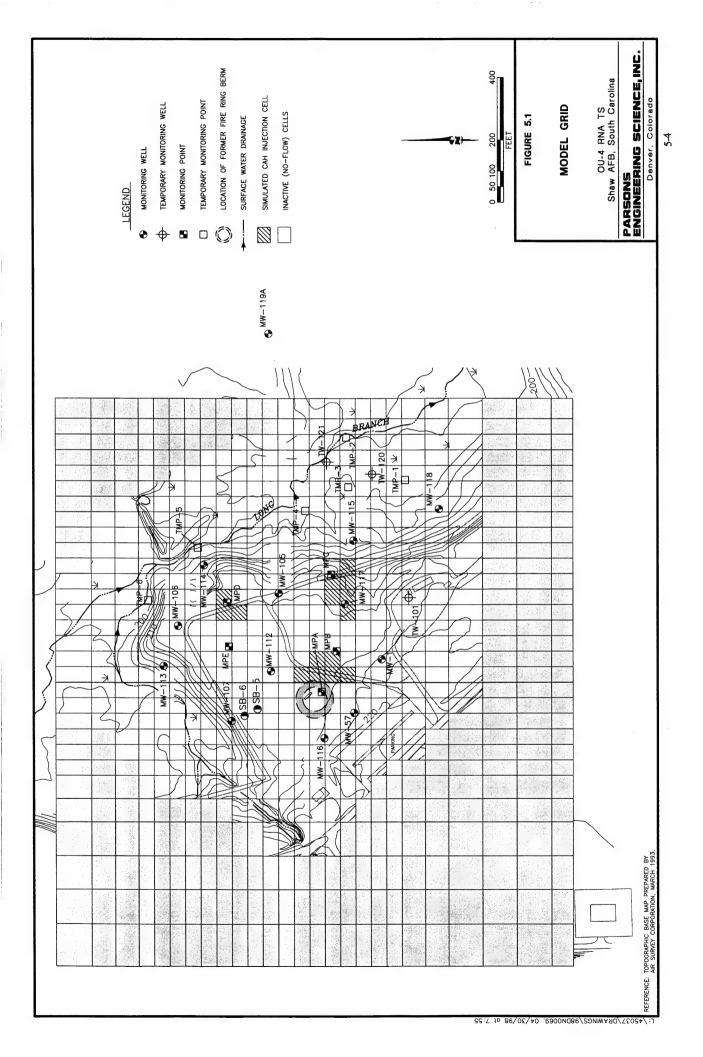
#### 5.3.1 Grid Design

The model domain for OU-4 is represented using one layer, with a 26- by 30-cell horizontal grid. Relatively small grid cells (50 feet by 50 feet) were used within the central portion of the contaminant plume to allow more accurate simulation of transport; the cell size was gradually increased with distance from the source area, with the largest cell measuring 140 feet wide by 110 feet long. The grid was oriented so that contaminant migration of the eastern plume lobe would run parallel to model rows. The grid thickness was assumed to span the elevations between 220 feet msl (general land surface elevation of the upper terrace) to 165 feet msl (approximate elevation of the geologic interface between medium to coarse and fine sands), or a model grid thickness of approximately 55 feet. The model grid covers an area of approximately 70 acres. The full extent of the model grid is indicated on Figure 5.1.

#### 5.3.2 Groundwater Flow Model

#### **5.3.2.1** Boundary Conditions

In defining the model domain, the area of interest must be separated from the surrounding system. Boundary conditions describe the interaction between the system being modeled and its surroundings or, for transport models, the loading of contaminant mass into the system. Boundary conditions are used to include the effects of the system outside the area being modeled with the system being modeled, while at the same time allowing the isolation of the desired model domain from the larger system. In effect, the boundaries of the model tell the area immediately inside the boundaries what to expect from the outside world. The solution of any differential equation requires specification of the conditions at the periphery of the system. Model boundaries are thus mathematical statements that specify the dependent variable (head



### TABLE 5.1 COMMON DESIGNATIONS FOR SEVERAL IMPORTANT BOUNDARY CONDITIONS<sup>a/</sup>

### SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

			General Mathem	atical Description
Boundary Condition	Boundary Type	Formal Name	Groundwater Flow	Contaminant Transport
Specified-Head or Specified- Concentration	Type One	Dirichlet	H = f(x, y, z, t)	C = f(x, y, z, t)
Specified Flux	Type Two	Neumann	$\frac{\partial H}{\partial n} = f(x, y, z, t)$	$\frac{\partial C}{\partial n} = f(x, y, z, t)$
Head- Dependent or Concentration- Dependent Flux	Type Three (mixed- boundary condition)	Cauchy	$\frac{\partial H}{\partial n} + cH = f(x, y, z, t)$	$\frac{\partial C}{\partial n} + cC = f(x, y, z, t)$

al Modified from Franke et al. (1987).

or contaminant concentration) or the flux (derivative of the head or contaminant concentration with respect to time) at the model grid boundaries.

Three types of boundary conditions generally are used to describe groundwater flow and solute transport. Boundary conditions are referred to as type one (Dirichlet), type two (Neumann), and type three (Cauchy). Table 5.1 summarizes boundary conditions for groundwater flow and solute transport.

In flow models, boundary conditions are ideally used to specify actual hydrogeologic boundaries to the system, such as streams, lakes, confining units, groundwater divides, or any geologic or anthropogenic feature that may bound a system. Also, the boundaries may be defined as areas where properties (e.g., flux) are known and can be defined. When using a numerical flow model, hydrologic boundaries such as constant-head features (e.g., lakes, etc.) or constant-flux features should, when possible, coincide with the perimeter of the model. In areas that lack obvious hydrologic boundaries, constant-head or constant-flux boundaries can be specified at the numerical model perimeter as long as the perimeter is far enough removed from the contaminant plume that transport calculations will not be affected by inaccuracies in the boundary conditions.

Specific-head cells were defined at the model boundaries. The specified heads were estimated by projecting heads from the groundwater flow maps. These constant-head cells were placed far enough from the simulated CAH plume to avoid potential boundary interferences. The MODFLOW river package was used to simulate Long Branch Creek and the adjoining tributary. Heads along Long Branch Creek were estimated to range from 200 feet msl at the northern boundary of the model domain to 191.5 feet msl at the southeastern boundary of the model domain. Heads of the adjoining tributary north of the site that feeds into Long Branch Creek were estimated

to range from 201.9 feet msl to 199.7 feet msl before connecting to Long Branch Creek. The base or lower boundary of the model is assumed to be no-flow and is set at 165 feet msl at the geologic interface of medium to coarse and fine sands. The upper model boundary is defined by the simulated water table surface.

### 5.3.2.2 Recharge and Evapotranspiration

Precipitation recharge is important at OU-4 because of high annual precipitation rates, sparsely vegetated terrain immediately surrounding the former fire-training pit, a relatively thin vadose zone (e.g., groundwater is at ground surface in the marshy terrain bordering Long Branch Creek), minimal surface paving, and permeable surface and subsurface geology. A recharge rate of 40 inches per year was assumed for non-forested areas at OU-4 [compared to an annual precipitation rate of 46 inches per year (Section 3)]. Because evapotranspiration is expected to increase in the forested areas at the site, a recharge rate of 12 inches per year was assumed for the forested areas east and north of the source. Appendix C shows the recharge map used for the model.

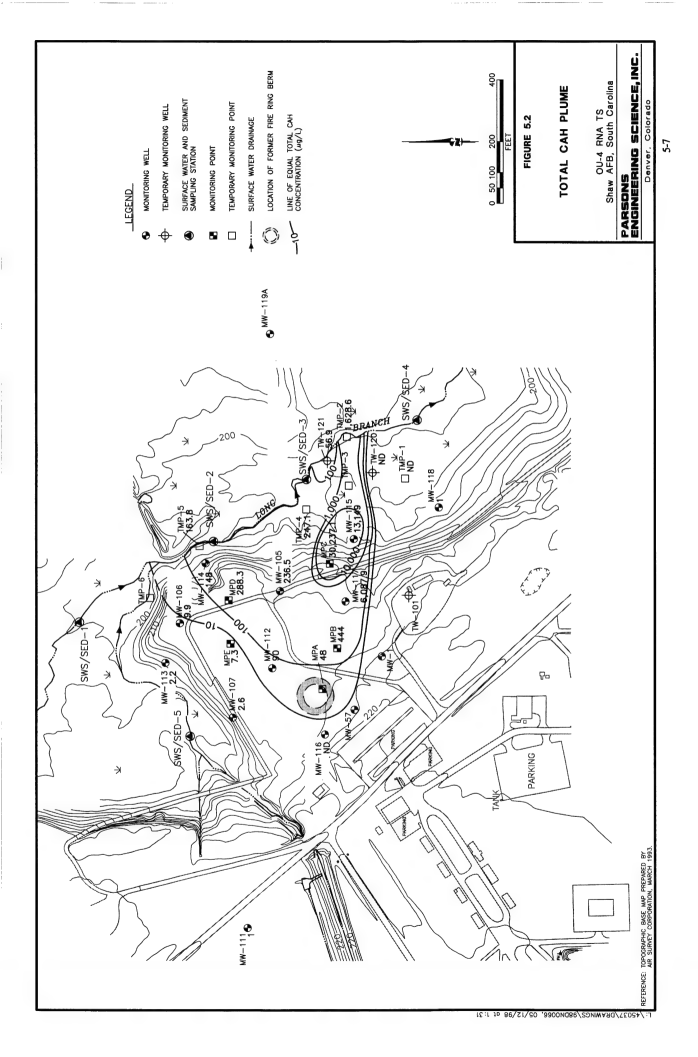
### 5.3.2.3 Aquifer Properties

As discussed in Section 3.3.2.3, the effective porosity is the percentage of a rock or sediment through which fluids can travel. A general value of 25 percent for effective porosity was defined for the model domain on the basis of the grain-size distribution observed in the shallow surficial aquifer (Spitz and Moreno, 1996).

Hydraulic conductivity values were calculated from field data obtained from 2 slug tests performed at monitoring wells MW-106 and MW-112, which are screened in the upper portion of the aquifer. Additional details on the slug tests are discussed in Section 3.3.2.2. It was assumed that the majority of the contaminant plume is migrating through the medium to coarse sands comprising the upper surface of the aquifer. An average hydraulic conductivity of 75 ft/day was initially defined for the model domain.

### 5.3.3 Contaminant Transport Model

The total dissolved CAH concentrations (combining chlorinated ethene and ethane concentrations obtained from May 1997 laboratory analytical results) for each monitoring well/point location were used as targets for model calibration. Table 4.6 presents dissolved CAH concentration data for May 1997, and Figure 5.2 shows the spatial distribution of the total dissolved CAH plume in May 1997. Predicting the natural attenuation of the total CAH plume is more appropriate than simulating the natural attenuation of a single CAH compound, such as 1,1,1-TCA. For instance, the compound 1,1-DCE results from both the reductive dehalogenation of TCE and from the dehydrohalogenation of 1,1,1-TCA. Therefore, all three compounds must be included to accurately simulate the fate and transport phenomenon of the chlorinated solvent plume. Furthermore, 1,1-DCA is the reductive dechlorination byproduct of 1,1,1-TCA and was present at a maximum concentration of 8,700 µg/L at monitoring well MW-115. 1,1-DCA currently does not have federally mandated MCLs; however, it contributes a significant amount of biodegradable chlorinated solvent mass to the groundwater system and was included in the model simulation.



### 5.3.3.1 Source

Transport models use boundary conditions to express the influence of contaminant sources such as nonaqueous-phase liquid (NAPL) bodies and dissolved mass entering through recharge, injection wells, surface water bodies, and leaking structures. Sources such as NAPL bodies may be represented as specified-concentration boundaries (limited by solubility constraints or observed maximum concentrations) or as specified-flux boundaries (for which the chemical loading rate must be known or estimated). However, in most cases, only the effects of the source are measured, not the source characteristics (Spitz and Moreno, 1996). The source must therefore be represented as a "black box" that produces appropriate contaminant concentrations or fluxes at selected points in the model. The source may be misrepresented under such a scenario, but there is often little choice in the matter. Estimating contaminant flux into groundwater from NAPL is difficult and is dependent upon several parameters, most of which cannot be measured (Feenstra and Guiguer, 1996; Abriola, 1996).

Rather than using various calculations to attempt to estimate CAH partitioning from NAPL into groundwater, the "black box" source approach was used. Experience modeling contaminated sites as part of the AFCEE Natural Attenuation Initiative has suggested that this is the best method for reproducing observed plumes. At the OU-4 site, the contaminant is assumed to enter groundwater in the study area through contact between groundwater and residual NAPL at or below the water table. Partitioning of CAHs from this source into groundwater was simulated by defining source area recharge concentrations in 15 model cells located at two known and one suspected source location at the site (Figure 5.1). The suspected source was located northeast of the fire training berm near monitoring point where a thin product sheen was detected. Low CAH concentrations upgradient of monitoring point MPD suggest that the increase in groundwater CAH concentrations at MPD were the result of contamination near this location and not from the fire training pit. The concentrations at all source locations were set at high concentrations to simulate the dissolution of current NAPL sources into the groundwater, and then adjusted (Section 5.4.2) to obtain the present observed effect.

### 5.3.3.2 Dispersivity

Published data summarized by Spitz and Moreno (1996) suggest that, as a rule of thumb, longitudinal dispersivity is approximately one-tenth the travel distance of the plume (from the source to the downgradient toe). For purposes of this model, dispersivity values were initially set at 50 feet which is approximately one-tenth the length of the east/southeast contaminant plume lobe where most of the groundwater contamination resides (Figure 5.2). Transverse dispersivity values were estimated as one-tenth of the longitudinal dispersivity values (Domenico and Schwartz, 1990).

### 5.3.3.3 Retardation

Retardation of CAHs relative to the advective velocity of the groundwater occurs when contaminant molecules are sorbed to organic carbon, silt, or clay in the aquifer matrix. Based on a measured TOC concentration of 0.00024 (Rust E&I, 1995), an assumed bulk density of 1.72 kilograms per liter (kg/L), and published values of the soil sorption coefficient (K<sub>oc</sub>) for DCE, DCA, and TCA (as listed in Montgomery *et* 

al., 1990) the coefficient of retardation for each contaminant was calculated. The results of these calculations are summarized in Table 5.2. The lower the assumed coefficient of retardation, the faster the CAH plume will migrate downgradient. Initially, the average calculated retardation coefficient of 1.12 was assigned to the model.

### 5.3.3.4 Biodegradation

As discussed in Section 4.5, first-order decay rates of 2.5 x 10<sup>-3</sup> day<sup>-1</sup> to 1.4 x 10<sup>-2</sup> day<sup>-1</sup> were calculated for CAH contamination using site-specific data. These rates were used to define a range of possible values for model input. An estimate of 5.5 x 10<sup>-3</sup> days<sup>-1</sup> was defined for the model domain. Also as noted in Section 4.5, reductive dechlorination of CAHs may be best represented using a second-order rate, but most common transport codes (including MT3D<sup>96</sup>) can incorporate only a first-order rate. This impacts model calibration and may constrain the usefulness of predictions, but at this time there is no feasible alternative for simulating CAH transport under the influence of biodegradation. Provided that conservative values are used for other input parameters and a sensitivity analysis is performed, simulations incorporating a first-order rate constant will still be useful and meaningful.

### 5.4 MODEL CALIBRATION

Model calibration is an important component in the development of any numerical groundwater model. Calibration of the flow model demonstrates that the model is capable of matching hydraulic conditions observed at the site; calibration of a contaminant transport model superimposed upon the calibrated flow model helps demonstrate that contaminant loading and transport conditions are being appropriately simulated. Model input and output are included in Appendix D.

### **5.4.1 Groundwater Flow Model**

Groundwater elevation data collected in May 1997 and presented on Figure 3.5 were used to calibrate the flow model. Water level elevation data from 11 monitoring wells were used to compare measured and simulated heads for calibration. The selected locations were TW-101, MW-105, MW-106, MW-107, MW-112, MW-113, MW-114, MW-115, MW-116, MW-117, and MW-118.

The numerical flow model was calibrated by altering hydraulic conductivity, constant-head elevations at the model boundaries, and the river package parameters in a trial-and-error fashion until simulated heads approximated observed field values within a prescribed accuracy. Figure 5.3 shows the calibrated water table. Calibrated model hydraulic conductivities ranged between 10 and 75 ft/day, with most cells having a hydraulic conductivity of 75 ft/day.

The root mean squared (RMS) error is commonly used to express the average difference between simulated and measured heads. RMS error is the average of the squared differences between measured and simulated heads, and can be expressed as:

### TABLE 5.2

# CALCULATION OF RETARDATION COEFFICIENTS (Foc = 0.00024) OU4 RNA TS

# SHAW AFB, SOUTH CAROLINA

			Fraction	Distribution	Distribution Distribution	Bulk	Bulk		
	Molecular	K %	Organic	Coefficient	Coefficient	Density	Density	Effective	Coefficient of
Compound	Weight	$(L/kg^{sl})$	Carbon b/	K <sub>d</sub> (L/kg)	$K_d$ (ft <sup>3</sup> /kg)	$(kg/L)^{c'}$	$(kg/ft^3)$	Porosity <sup>d/</sup>	Retardation
PCE	167.85	263	0.00024	0.063	0.00223	1.72	48.70	0.25	1.43
TCE	131.39	107	0.00024	0.026	0.00091	1.72	48.70	0.25	1.18
1,1-DCE	96.94	65	0.00024	0.016	0.00055	1.72	48.70	0.25	1.11
cis-1,2-DCE	96.94	65	0.00024	0.016	0.00055	1.72	48.70	0.25	1.11
trans-1,2-DCE	96.94	59	0.00024	0.014	0.00050	1.72	48.70	0.25	1.10
VC	62.5	2	0.00024	0.001	0.00002	1.72	48.70	0.25	1.00
1,1,1-TCA	133.4	126	0.00024	0.030	0.00106	1.72	48.70	0.25	1.21
1,1-DCA	98.86	30	0.00024	0.007	0.00026	1.72	48.70	0.25	1.05
Chlorobenzene	112.56	126	0.00024	0.030	0.00107	1.72	48.70	0.25	1.21
1,2-Dichlorobenzene	147	186	0.00024	0.045	0.00158	1.72	48.70	0.25	1.31
1,3-Dichlorobenzene	147	170	0.00024	0.041	0.00144	1.72	48.70	0.25	1.28
1,4-Dichlorobenzene	147	159	0.00024	0.038	0.00134	1.72	48.70	0.25	1.26
Benzene	84	42	0.00024	0.019	0.00067	1.72	48.70	0.25	1.13
Toluene	86	190	0.00024	0.046	0.00161	1.72	48.70	0.25	1.31
Ethylbenzene	112	468	0.00024	0.112	0.00397	1.72	48.70	0.25	1.77
Xylenes	112	394	0.00024	0.095	0.00334	1.72	48.70	0.25	1.65

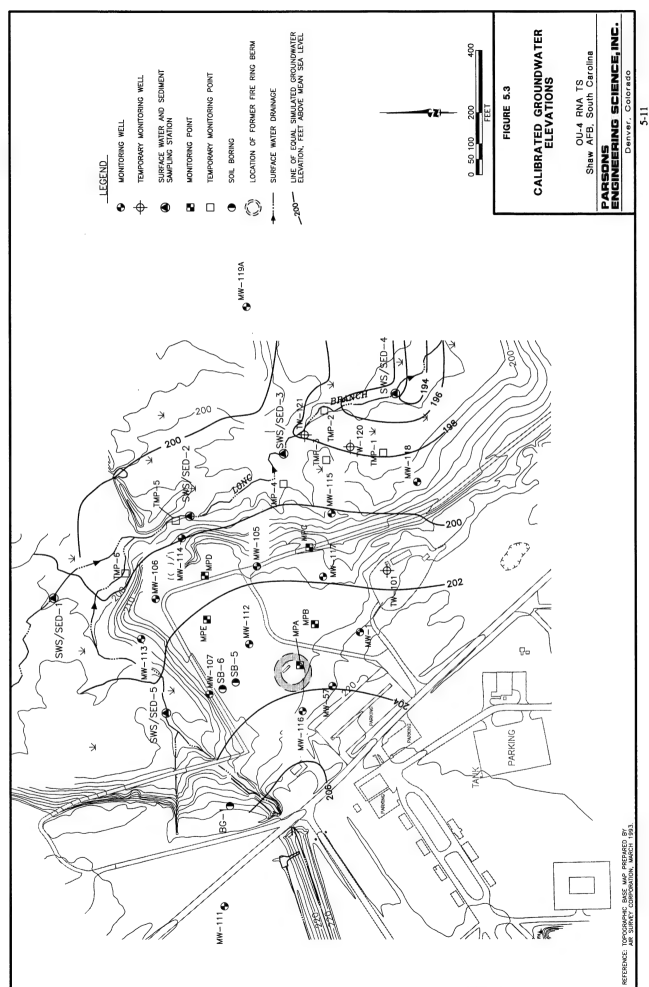
### NOTES:

<sup>&</sup>lt;sup>2</sup> From technical protocol (Wiedemeier et al., 1996), Knox et al. (1993), and Montgomery et al., (1990).

<sup>&</sup>lt;sup>b/</sup> FOC value measured from soil samples at the site (Rust, 1995).

Assumes an aquifer grain density of 2.65 (quartz sand) and a volumetric porosity of 35%.

<sup>&</sup>lt;sup>d'</sup> An effective porosity of 0.25 was estimated in the RFI (Parsons ES, 1997).



RMS = 
$$\left[\frac{1}{n}\sum_{i=1}^{n}(h_{m}-h_{s})_{i}^{2}\right]^{0.5}$$

where:

n = the number of points where heads are being compared,

 $h_m$  = measured head value, and

 $h_s = \text{simulated head value}.$ 

The RMS error between observed and calibrated values at the 11 comparison points was 0.28 feet, which corresponds to a calibration error of 6.4 percent (water levels dropped approximately 4.51 feet over the portion of the model domain containing the monitoring wells/points listed above). In solving the groundwater flow equation, Visual MODFLOW establishes the water table surface and calculates an overall hydraulic balance that accounts for the numerical difference between flux into and out of the system. The hydraulic mass balance for the calibrated model was adequate to accomplish the objectives of this modeling effort, with a 0-percent discrepancy.

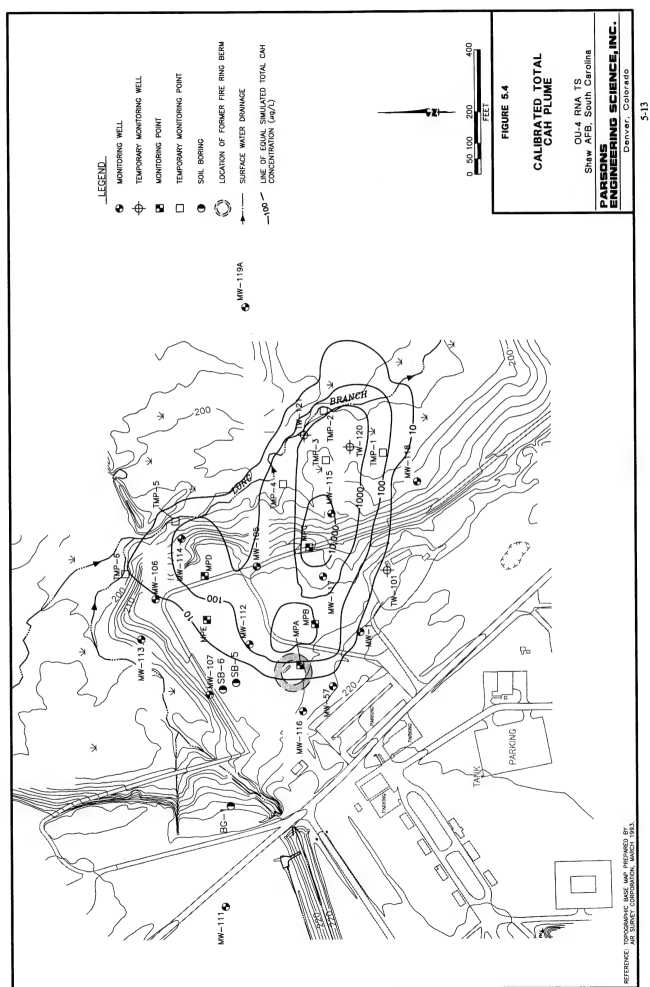
### 5.4.2 Plume Calibration

After calibration of the flow model, the numerical solute transport model was calibrated by altering the contaminant specified-flux rate in the source cells and contaminant transport parameters in a trial-and-error fashion until the simulated plume approximated observed field values. The transport parameters varied during plume calibration were the source concentrations and the plume dispersivity. Because the original estimates for these parameters resulted in a calculated plume that reasonably resembled the original plume, these parameters were not varied greatly.

The dissolved CAH concentrations obtained from April 1997 laboratory analytical data from 23 monitoring wells/points sampled by Parsons ES were used to calibrate the contaminant transport model. The calibration points were MW-105, MW-106, MW-107, MW-112, MW-113, MW-114, MW-115, MW-116, MW-117, MW-118, TW-101, TW-120, TW-121, MPA, MPB, MPC, MPD, MPE, TMP-1, TMP-2, TMP-3, TMP-4, and TMP-5.

The longitudinal dispersivity of 50 feet produced a reasonable simulated plume but was decreased to 10 feet in the source area because dispersivity is scale-dependent. This is the value used to produce the calibrated plume depicted on Figure 5.4.

The calibrated plume calculated by the model (Figure 5.4) is similar, but not identical, to the observed May 1997 plume (Figure 5.2). Simulated CAH concentrations in the three simulated source areas are slightly higher than observed March 1997 concentrations. The simulated total CAH concentration at temporary monitoring point MPC is  $30,260~\mu g/L$ , whereas the concentrations observed at that well was  $30,237~\mu g/L$ . CAH concentrations simulated at this location were the largest in the



model. Simulated CAH concentrations at the other two source areas were 57  $\mu$ g/L (MPA within former fire training berm) and 307  $\mu$ g/L (MPD where groundwater sheen detected) compared to actual concentrations of 48 and 288  $\mu$ g/L, respectively. Simulated CAH concentrations at monitoring points adjacent to the creek varied. For instance, the measured CAH concentrations at TW-121 (edge of east/southeast plume lobe) and TMP-5 (downgradient edge of northeast plume lobe) were 424  $\mu$ g/L and 75  $\mu$ g/L versus observed concentrations of 57 and 164  $\mu$ g/L, respectively. In general, simulated CAH concentrations at the plume toe exceeded observed concentrations.

Higher simulated concentrations in the source area indicate that an adequate amount of mass is being introduced to the modeled system. Higher simulated concentrations midway through the plume shows that the model conservatively accommodates more mass than there is reason to believe is present. Similarly, higher simulated groundwater concentrations near the edge of the creek probably will exceed actual groundwater concentrations. Therefore, the conservative features of the model by maintaining simulated groundwater concentrations above observed groundwater concentrations dictates that the model predicts the worst possible groundwater impacts at the site.

### 5.5 SENSITIVITY ANALYSIS

The purpose of the sensitivity analysis is to determine the effect of varying model input parameters on model output. The sensitivity analysis was conducted by varying hydraulic conductivity, the coefficient of retardation, the first-order decay rate, dispersivity, and the CAH injection rate.

To perform the sensitivity analyses, the calibrated model was adjusted by systematically changing the aforementioned parameters individually, and then comparing the new simulations to the results of the calibrated model. The models were run for a 2-year period, just as the calibrated model was, so that the independent effect of each variable could be assessed. Eight sensitivity runs of the calibrated model were made, with the following variations:

- A. Hydraulic conductivity uniformly increased by a factor of 2;
- B. Hydraulic conductivity uniformly decreased by a factor of 2;
- C. Dispersivity uniformly increased by a factor of 2;
- D. Dispersivity uniformly decreased by a factor of 2;
- E. Coefficient of retardation uniformly increased by 50 percent;
- F. Coefficient of retardation uniformly decreased by 50 percent;
- G. First order decay rate increased by an order of magnitude; and
- H. First order decay rate decreased by an order of magnitude.

The results of the sensitivity analyses are discussed in the following subsections and primarily address sensitivity variations in the east/southeast plume lobe, which has the highest total CAH concentrations at the site. As described in the following paragraphs, the parameter modifications listed above generally resulted in significant changes in the resulting plumes, with the retardation modifications having the smallest effect.

### 5.5.1 Sensitivity to Variations in Hydraulic Conductivity

Hydraulic conductivity is an important aquifer characteristic that represents the ability of the water-bearing strata to transmit groundwater. An accurate estimate of hydraulic conductivity is important to help quantify advective groundwater flow velocities and to define the flushing potential of the aquifer. As a result, models used to estimate contaminant transport are particularly sensitive to variations in hydraulic conductivity. Lower values of hydraulic conductivity result in a slower-moving plume. Higher values of hydraulic conductivity result in a faster-moving plume. The effects of varying hydraulic conductivity are summarized in Table 5.3. Uniformly increasing the hydraulic conductivity by a factor of two decreased the lateral dispersion of the plume to cause a slight shrinking in plume size. Simulated total CAH concentrations in the source areas decreased as simulated throughput through the source area was increased. For instance, concentrations at monitoring well MPC decreased from 30,261 µg/L to 18,102 µg/L with increased hydraulic conductivity. Simulated total CAH concentrations were observed to increase near Long Branch Creek due to higher transport velocities and decreased residence times for biodegradation. In contrast, decreasing the hydraulic conductivity by a factor of two slowed overall plume migration, and caused the contaminant mass to be concentrated within a smaller area. As a result, CAH levels in the vicinity of source areas increased (monitoring point MPC increased from 30,261 µg/L to 46,732 µg/L) and the simulated CAH concentrations near Long Branch Creek uniformly decreased (monitoring wells OBS-A and TMP-5 decreased by approximately 1,215 and 37 µg/L, respectively).

### 5.5.2 Sensitivity to Variations in Dispersivity

The effects of varying dispersivity are summarized in Table 5.3. Both longitudinal and transverse dispersivity were varied for this analysis, as the ratio of the two values was kept constant at 0.1. Increasing the dispersivity by a factor of two resulted in relatively lower concentrations in the source area. The simulated concentration at source area monitoring point locations MPC was 27,999  $\mu$ g/L, compared to a calibrated concentration of 30,261  $\mu$ g/L. Corresponding decreases were observed downgradient from the source area, causing the simulated concentration at well OBS-A to decrease from 2,103  $\mu$ g/L in the calibrated model to 1,929  $\mu$ g/L.

Decreasing the dispersivity by a factor of two produced a plume with slightly higher concentrations in the source area. The simulated concentration at monitoring point MPC was 31,538  $\mu$ g/L, compared to the calibrated model concentration of 30,261  $\mu$ g/L. The CAH concentrations in the downgradient portion of the plume also were higher. The simulated concentration at observation well OBS-A was 2,208  $\mu$ g/L, compared to the observed concentration of 2,103  $\mu$ g/L at that well.

TABLE 5.3
SENSITIVITY ANALYSIS RESULTS
OU4 RNA TS

# SHAW AFB, SOUTH CAROLINA

	Measured Total	Calibrated Total								
	CAH Plume	CAH Plume					Retardation	Retardation	Decay Rate Decay Rate	Decay Rate
Well No.	Concentration	Concentration					Increased	Decreased	by	Decreased
	May '97 $(\mu g/L)^{a'}$	$(\mu g/L)$	K x2 <sup>b</sup> /	K/2	$\alpha \times 2^{c'}$	$\alpha/2$	by 50%	by 50%	by 10	by 10
East/Sout	East/Southeast Plume Lobe									
MPA	48	57	36	11	150	18	25	58	7054	68
MPB	444	463	319	292	524	451	455	466	28	825
MPC	30237	30261	18012	4673	2799	31538	30169	30473	10577	39310
MW-117	8809	6856	4267	8566	7528	6390	6812	9069	1851	9862
MW-115	13149	13321	10545	13238	12366	13872	13093	13521	422	29687
OBS-A	1629 <sup>d/</sup>	2103	2752	888	1929	2208	2010	2172	0	11812
Northeast	Northeast Plume Lobe									
MPD	288	304	15	416	305	301	301	307	55	571
MW-114	148	169	164	129	164	171	165	173	2	593
TMP-5	164	75	82	38	78	73	72	78	0	391

a/  $\mu g/L = \text{micrograms per liter.}$ 

b/ K = hydraulic conductivity.

 $c/\alpha = dispersivity.$ 

d/ The measured value for monitoring point TMP-2 was used for observation well OBS-A.

Note: The adjustments made to the calibrated model during the various sensivity analysis runs are more fully described in Section 5.5.

### 5.5.3 Sensitivity to Variations in the Coefficient of Retardation

The effects of varying the coefficient of retardation are summarized in Table 5.3. Uniformly increasing the coefficient of retardation by 50 percent slightly limited the downgradient migration of the plume. The simulated total CAH molar concentration in the vicinity of source area monitoring point MPC was 30,169 µg/L, compared to the calibrated concentration of 30,261 µg/L. A small decrease in the downgradient concentrations of the total CAH plume occurred, as evidenced by a simulated CAH concentration of 2,010 µg/L at observation well OBS-A, where the calibrated CAH concentration was 2.103 ug/L. In contrast, decreasing the coefficient of retardation by 50 percent slightly increased plume migration. As a result, the simulated CAH concentration at source area monitoring point MPC increased by approximately 200 ug/L as source areas upgradient from MPC released more contaminant mass downgradient. Concentrations downgradient from the source area also increased (CAH concentrations changed from 2,103 to 2,172 µg/L at observation well OBS-A). Overall, varying the coefficient of retardation within a reasonable range has a limited effect on contaminant transport.

### 5.5.4 Sensitivity to Variations in the Anaerobic Decay Rate Constant

The effects of varying the first-order CAH decay rate are summarized in Table 5.3. Increasing this parameter by an order of magnitude results in more rapid degradation of dissolved contaminants. The resulting simulated CAH concentrations at source area well MPC and at downgradient observation well OBS-A were lower than in the calibrated model (10,577  $\mu$ g/L and 0  $\mu$ g/L, respectively). Increasing the degradation rate by an order of magnitude results in a much smaller plume. In contrast, decreasing this parameter by an order of magnitude causes a large increase in CAH concentrations throughout the entire plume. For instance, groundwater CAH concentrations at monitoring point MPC and observation well OBS-A increased to 39,310 and 11,812  $\mu$ g/L, respectively.

### 5.5.6 Summary of Sensitivity Analysis Results

The results of the sensitivity analysis suggest that the calibrated model depicted in Figure 5.4 is reasonable. Varying the model parameters within the prescribed ranges generally caused the extent and magnitude of the dissolved plume to differ substantially from observed site conditions.

### **SECTION 6**

### COMPARATIVE ANALYSIS OF REMEDIAL ALTERNATIVES

This section presents the development and comparative analysis of two groundwater remedial alternatives for OU-4 at Shaw AFB. The intent of this evaluation is to determine if RNA is an appropriate and cost-effective remedial approach to consider when developing final remedial strategies for the CAHs dissolved in site groundwater, especially when combined with other innovative and conventional remedial technologies. A groundwater extraction and treatment system consisting of at least two extraction wells will be installed as an interim measure (IM) during the spring of 1998 near wells MW-117 and MW-112. The intent of this system is to mitigate contaminant plume migration until the option of monitored natural attenuation is evaluated, largely through the results of this TS. The predicted effects of an interim, limited-term (two year) groundwater extraction and treatment system, combined with natural attenuation, The modeling effort did not include predictions to are described in this section. determine the future extent and concentration of the BTEX plume at OU-4 due to lower BTEX concentrations relative to CAH concentrations, the role of BTEX as electron donors for CAH degradation, and the fact that detectable concentrations of BTEX are not present in surface water.

Section 6.1 presents the criteria used to evaluate groundwater remedial alternatives. Section 6.2 discusses the development of remedial alternatives considered as part of this demonstration project. Section 6.3 provides a brief description of each of these remedial alternatives. Section 6.4 provides a more detailed analysis, including modeling results, of the remedial alternatives using the defined evaluation criteria. The results of this evaluation process are summarized in Section 6.5.

### 6.1 REMEDIAL ALTERNATIVE EVALUATION CRITERIA

The evaluation criteria used to identify appropriate remedial alternatives for shallow groundwater contamination at the site were adapted from those recommended by USEPA (1988) for selecting remedies for Superfund sites (OSWER Directive 9355.3-01). These criteria include (1) long-term effectiveness and permanence, (2) technical and administrative implementability, and (3) relative cost. The following sections briefly describe the scope and purpose of each evaluation criterion. This report focuses on the potential use of RNA to reduce CAH concentrations within the shallow groundwater to levels that meet regulatory standards intended to be protective of human health and the environment.

### **6.1.1 Long-Term Effectiveness and Permanence**

Each remedial approach or remedial alternative (which can be a combination of remedial approaches such as RNA and institutional controls) was analyzed to determine how effectively it will minimize groundwater plume expansion so that applicable groundwater quality standards can be achieved at a downgradient POC. The expected remedial effectiveness based on case histories from other sites with similar conditions The ability to minimize potential impacts on also is evaluated where feasible. surrounding areas, facilities, and operations is considered. Also, the ability of each remedial alternative to protect both current and potential future receptors from potential exposures associated with site-related contamination in shallow groundwater is qualitatively assessed by conservatively evaluating the potential for completion of exposure pathways involving groundwater, either now or in the future. This evaluation criterion also includes permanence and the ability to reduce contaminant mass, toxicity, and volume. Time to implementation and estimated time until protection is achieved are described. Long-term reliability for providing continued protection, including an assessment of potential for failure of the technology and the potential threats resulting from such a failure, also is evaluated.

### **6.1.2** Implementability

The technical implementation of each remedial technology/approach or remedial alternative was evaluated in terms of technical feasibility and availability. Potential logistical shortcomings and difficulties in construction, operations, and monitoring are presented and weighed against perceived benefits. Requirements for any post-implementation site controls such as LTM and land or groundwater use restrictions are described. Administrative feasibility in terms of public acceptance and the ability to obtain necessary approvals is discussed.

### 6.1.3 Cost

The total cost (present worth) of each remedial alternative was estimated for relative comparison following USEPA (1993) guidance. An estimate of capital costs, and operations and post-implementation costs for site monitoring and controls is included. An annual adjustment (discount) factor of 7 percent was assumed in present-worth calculations (USEPA, 1993).

### 6.2 FACTORS INFLUENCING ALTERNATIVES DEVELOPMENT

Several factors were considered during the identification and screening of remedial technologies for addressing shallow groundwater contamination at the site. Factors considered included the objectives of the natural attenuation demonstration program; contaminant, groundwater, and soil properties; present and future land uses; and potential receptor exposure pathways. The following section briefly describes each of these factors and how they were used to narrow the list of potentially applicable remedial technologies to the final remedial alternatives considered for the site.

### 6.2.1 Program Objectives

The intent of the RNA demonstration program sponsored by AFCEE is to develop a systematic process for scientifically investigating and documenting naturally occurring subsurface attenuation processes that can be factored into overall site remediation plans. The objective of this program is to provide solid evidence of RNA of dissolved CAHs so that this information can be used to develop an effective groundwater remediation strategy. A secondary goal of this multi-site initiative is to provide a series of regional case studies that demonstrate that natural processes of contaminant degradation can often reduce contaminant concentrations in groundwater to below acceptable cleanup standards before completion of potential exposure pathways.

Because the objective of this program is to study natural processes in the saturated zone rather than all contaminated media (vadose zone soil, soil gas, etc.), remedial approaches and technologies have been evaluated based primarily on their potential impact on shallow groundwater and phreatic soils. Technologies that can reduce vadose zone contamination and partitioning of contaminants into groundwater also have been evaluated. Many of the source removal technologies evaluated in this section also will reduce soil and soil gas contamination, but it is important to emphasize that the remedial alternatives developed in this document are not necessarily intended to remediate all contaminated media.

Additional program objectives set forth by AFCEE include cost effectiveness and minimization of waste. Approaches and technologies that may meet these criteria include institutional controls, soil vapor extraction, air sparging, and RNA. Soil excavation, slurry walls, sheet piling, carbon adsorption, ex situ biological or chemical treatment, groundwater pump and treat, and onsite/offsite disposal generally are not attractive technology candidates under this program.

### **6.2.2 Contaminant Properties**

The site-related chlorinated solvent contaminants evaluated as part of this TS include 1,1,1-TCA, 1,1-DCA, PCE, TCE, cis- and trans-1,2-DCE, 1,1-DCE, VC, DCB, and CB. Three contaminant sources areas have been identified; sources include LNAPL floating on the water table and/or residual contamination adsorbed to soils. Two of the source areas contribute relatively little to groundwater CAH contamination, whereas the third source area appears to cause concentrated CAH contamination in the east/southeast plume lobe (Section 4). The physiochemical characteristics of the individual CAH compounds within these plumes will greatly influence the effectiveness and selection of a remedial technology.

Several of the chlorinated solvents present at OU-4 were not evaluated in the selection of remedial alternatives for the site. The compounds DCB and CB were present below USEPA MCLs and were omitted from this evaluation. The remaining chlorinated solvents detected in site groundwater include 1,1,1-TCA, 1,1-DCA, 1,2-DCA, PCE, TCE, cis- and trans-1,2-DCE, 1,1-DCE, and VC. Most of these compounds were detected at relatively low concentrations and/or were detected at a single monitoring well/point location (e.g., trans-1,2-DCE and 1,2-DCA), and are not considered a primary groundwater threat. The compounds 1,1,1-TCA, cis-1,2-DCE,

and 1,1-DCA represent approximately 94 percent of the groundwater CAH plume mass.

1,1,1-TCA was used as the target remediation compound at OU-4. The summary of biodegradation rates in Table 4.9 indicates that the transformation rates of 1,1,1-TCA to daughter products (1,1,1-TCA comprises approximately 70 percent of all dissolved chlorinated ethane mass) versus the transformation rates of *cis*-1,2-DCE to VC (*cis*-1,2-DCE also comprises approximately 70 percent of all dissolved chlorinated ethene mass) were nearly identical. However, the maximum *cis*-1,2-DCE concentration was detected at the same location as the maximum 1,1,1-TCA concentration, but at a concentration approximately five times lower than 1,1,1-TCA. The relatively high 1,1,1-TCA concentrations support the use of this compound as a surrogate for all CAH compounds at this site during the remedial alternatives evaluation process. The selection of 1,1,1-TCA in this evaluation also is consistent with the selection of 1,1,1-TCA as the target remedial compound for interim measures being conducted at the site by IT Corporation (SCDHEC, 1997).

All of the CAHs detected at OU-4 are generally volatile, relatively soluble in water, and do not adsorb strongly to soil particles. These characteristics allow the CAHs to leach readily from DNAPL and contaminated soil into groundwater, and to migrate as dissolved contamination (Lyman et al., 1992). Many CAHs are susceptible to in situ degradation by both biotic and abiotic mechanisms under certain geochemical conditions.

1,1,1-TCA, the selected surrogate compound, is a volatile compound with a vapor pressure of 124 millimeters of mercury (mm Hg) at 25°C (Neely and Blau, 1985). Henry's Law Constants reported for 1,1,1-TCA range from 0.013 to 0.023 atmospherecubic meters per mole (atm-m³/mol) at 25°C (Ashworth *et al.*, 1988; Dewulf *et al.*, 1995; Hine and Mookerjee, 1975; and Hunter-Smith *et al.*, 1983). The solubility of pure 1,1,1-TCA in water at 25°C has been reported to range from 347 to 1,334 mg/L (Howard, 1990; Neely and Blau, 1985). 1,1,1-TCA biodegradation products vary according to the prevailing groundwater geochemistry, and are described in Section 4.2.

On the basis of these physiochemical characteristics, natural attenuation, soil vapor extraction, air sparging, groundwater extraction, and air stripping technologies could all be effective options for collecting, destroying, and/or treating CAHs at OU-4. In addition, *in situ* degradation of CAHs via permeable reaction walls (e.g., iron filings trenches) is a developing technology that holds promise for treatment of CAHs dissolved in groundwater.

### **6.2.3 Site-Specific Conditions**

Two general categories of site-specific characteristics were considered when identifying remedial technologies for comparative evaluation as part of this demonstration project. The first category included physical characteristics such as groundwater depth, gradient, flow direction, and soil type, which influence the types of remedial technologies most appropriate for the site. The second category involved assumptions about future land uses and potential receptor exposure pathways. Each of

these site-specific characteristics have influenced the selection of remedial alternatives included in the comparative evaluation.

### 6.2.3.1 Groundwater and Soil Characteristics

Site geology and hydrogeology will have a profound effect on the transport of contaminants and the effectiveness and scope of required remedial technologies at a given site. Hydraulic conductivity is perhaps the most important aquifer parameter governing groundwater flow and contaminant transport in the subsurface. The velocity of the groundwater and dissolved contamination is directly related to the hydraulic conductivity of the saturated zone. Slug tests conducted in OU-4 indicate a moderate to high conductivity within the Duplin Formation terrace deposits, which primarily consist of unconsolidated, well- to poorly-graded sands. Estimated conductivity values for these deposits ranged from 18 ft/day (based on a pump test) to 81 ft/day (based on slug The shallow groundwater migrates to the northeast, east, and southeast, increasing the extent of contamination from the source area to Long Branch Creek (i.e., the plume has expanded). However, the CAH concentrations have decreased within the aquifer with distance from the source areas through dilution and biodegradation. Groundwater migration from the source areas to Long Branch Creek takes two years given the estimated groundwater velocities at OU-4 (Section 3). The combination of rapid groundwater velocities and discontinued fire training operations in 1969 indicate that steady-state or shrinking plume conditions currently are achieved at OU-4.

Although higher hydraulic conductivities have resulted in plume expansion and migration toward the bounding creek, this same characteristic also will enhance the effectiveness of remedial technologies such as groundwater and soil vapor extraction and air sparging. For example, it should be less expensive and time-consuming to capture and treat the contaminant plume using a network of extraction wells in the highly conductive soils. Contaminant recovery also may be maximized when contaminants are not significantly sorbed to and retarded by phreatic soil, as in the predominantly sandy soils at the site. The effectiveness of air sparging also may be increased in highly conductive aquifers because of reduced entry pressures and increased radius of influence. Greater hydraulic conductivity also would increase the amount of contaminant mass traveling through an air sparging network within a given time period.

To satisfy the requirements of 1,1,1-TCA biodegradation, the aquifer must provide an adequate and available carbon or energy source, relatively reducing conditions, suitably low concentrations of competing electron acceptors (e.g., DO), and proper ranges of pH and temperature. Data collected as part of the field work phase of this demonstration project (Sections 3 and 4) indicate that these conditions are present in the source areas of the groundwater plume; thereby contributing to the degradation of highly chlorinated CAHs through reductive dechlorination and additional abiotic degradation of 1,1,1-TCA through dehydrohalogenation. Indigenous microorganisms capable of degrading CAHs tend to be common in most soil environments, and appear to be active at OU-4. Indigenous microorganisms have a distinct advantage over microorganisms injected into the subsurface to enhance biodegradation because indigenous microorganisms are well adapted to the physical and chemical conditions of the subsurface in which they reside (Goldstein et al., 1985). Therefore, microbe addition is not considered a viable remedial technology for this site.

### **6.2.3.2** Potential Exposure Pathways

A pathways analysis identifies the human and ecological receptors that could potentially come into contact with site-related contamination and the contaminant migration pathways through which these receptors might be exposed. To have a completed exposure pathway, there must be a source of contamination, mechanism(s) of release, a pathway of transport to an exposure point, an exposure point, and a receptor. If any of these elements do not exist, the exposure pathway is considered incomplete, and receptors will not come into contact with site-related contamination. Evaluation of the potential long-term effectiveness of any remedial technology or remedial alternative as part of this demonstration project includes determining if the approach will be sufficient and adequate to minimize plume expansion so that potential receptor exposure pathways involving shallow groundwater contaminants are incomplete.

Assumptions about current and future land uses at a site form the basis for identifying potential receptors, exposure pathways, reasonable exposure scenarios, and appropriate remediation goals. USEPA (1988) advises that the land use associated with the highest (most conservative) potential level of exposure and risk that can reasonably be expected to occur should be used to guide the identification of potential exposure pathways and to determine the level to which the site must be remediated.

Most of OU-4 is a grassland/shrub habitat generally devoid of large vegetation. The site becomes more heavily vegetated with bottomland forest adjacent to Long Branch Creek. The former fire-training berm itself lacks any vegetation. The site is undeveloped with the exception of a picnic area located approximately 200 feet west of the former fire training berm. Therefore, the primary use of OU-4 is for recreational activities, and it is estimated that the picnic area is used 2 to 3 times per month during the spring to early fall months (Rust E&I, 1995).

Rural/residential land begins within 1,000 feet north and east of OU-4 and primarily is used for agricultural purposes. The areas south and west of the site are occupied by Base facilities. A Munitions Flight unit is located approximately 350 feet southwest of the site and a equipment storage area is located approximately 400 feet southeast. The water from Long Branch Creek discharges to a privately owned pond called Booths Pond, located approximately 1,000 feet south/southeast from the site. Groundwater sampling results indicate that the CAH plume may currently be impacting water quality in Long Branch Creek. However, as expressed in the baseline risk assessment (BRA) (Rust E&I, 1995), contaminants from other sources along Long Branch Creek also could be transported to Booths Pond and the pond was, therefore, not considered in the BRA.

Under reasonable current land use assumptions, potential receptors include Base worker populations, Base residents, residents living southeast of the Base, trespassers who enter the site, and recreators in Booths Pond or Long Branch Creek. The nearest employees to the site on a daily basis work at the Munitions Flight Unit. However, Munitions Flight Unit personnel do not come into direct contact with the site on a daily basis, although they utilize the parking area adjacent to the site. Potential contact with site contaminants could come from adults and children wading in Long Branch Creek while hunting, fishing, or playing. The ingestion of fish caught from the creek or Booths Pond would be another potential exposure route. According to Rust (1995),

there are no potable water supply wells at OU-4. The nearest supply wells (Base Wells 8 and 10) are located in the Munitions Flight Unit area and have been abandoned. Exposure pathways involving other environmental media such as vadose zone soils and soil gas in the source area were not considered as part of this project, but should be considered in overall site remediation decisions.

Assumptions about hypothetical future land uses also must be made to ensure that the remedial technology or alternative considered for shallow groundwater at the site is adequate and sufficient to provide long-term protection. The future use of OU-4 and the surrounding area is projected to be unchanged from the current uses described above. However, the possibility of Shaw AFB closing in the future exists, and land use could change. As part of the BRA (Rust E&I, 1995), future land use has conservatively been assumed to include on-site residential. Future receptor exposure to groundwater could occur through the use of an on-site well screened in the shallow aquifer and subsequent groundwater ingestion, dermal absorption while bathing, or inhalation of vapors during showering. Exposure to contaminants in a residential exposure scenario also could arise through absorption and ingestion during recreational contact with surface waters of Long Branch Creek or Booths Pond.

In summary, the use of RNA at this site will require that the source area be maintained as industrial or open-space (undeveloped) property and that restrictions on shallow groundwater use be enforced in areas downgradient from the site until natural attenuation reduces contaminants to concentrations that meet regulatory standards. Although shallow groundwater is in hydraulic communication with Long Branch Creek, contaminant concentrations detected in surface water were low (maximum CAH concentration was 2.3 μg/L cis-1,2-DCE at SWS-3). The substantial estimated flow volume of Long Branch Creek [>1,000 gallons per minute (gpm) (Rust E&I, 1995)], aerobic degradation processes, photooxidation, and volatilization are expected to dilute and degrade any CAH compounds entering the creek. If source removal technologies such as soil vapor extraction or groundwater pump and treat are implemented, they will have some impact on the short- and long-term land use options and also will require some level of institutional control and worker protection during remediation.

### 6.2.3.3 Remediation Goals for Shallow Groundwater

Potentially applicable state and federal water quality criteria for 1,1,1-TCA are summarized in Table 6.1.

Use of RNA assumes that compliance with promulgated drinking water standards is not necessary if site-related contamination does not pose an immediate threat to human health or the environment (i.e., exposure pathways are incomplete). Thus, the magnitude of required remediation in areas that can and will be placed under institutional control is different from the remediation that is required in areas that may be available for unrestricted groundwater use. The primary remedial action objective (RAO) for groundwater at and downgradient from OU-4 is limited plume expansion to prevent exposure of site receptors to concentrations of CAHs in groundwater and surface water at levels that exceed regulatory standards or guidelines. This means that viable remedial alternatives must be able to achieve concentrations that minimize plume migration and/or expansion. Institutional controls are likely to be a necessary component of any groundwater remediation strategy for this site. The required

### TABLE 6.1 REMEDIAL ACTION OBJECTIVES FOR GROUNDWATER AND SURFACE WATER

### OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

Compound	USEPA MCL³' (µg/L)b'	Interim Measure Remedial Goal <sup>c</sup> (µg/L)	Federal Ambient Surface Water Quality, Ingestion of Organisms d' (µg/L)
1,1,1-Trichloroethane	200	1,000	170,000

- <sup>a'</sup> USEPA MCL = US Environmental Protection Agency (1996) Maximum Contaminant Level.
- $\mu g/L = micrograms per liter.$
- South Carolina Department of Health and Environmental Control (SCDHEC, 1997).
- d' USEPA (1991). Note: This standard relates to the possible ingestion of aquatic organisms in Long Branch Creek.

duration of these institutional controls may vary depending on the effectiveness of the selected remedial technology at reducing contaminant mass and concentration in the groundwater, and the amount of contaminated groundwater discharging to surface water.

### 6.2.4 Summary of Remedial Technology Screening

Several remedial technologies were identified and screened for use in treating the Table 6.2 identifies the initial remedial technologies groundwater at the site. considered as part of this demonstration and those retained for detailed comparative Screening was conducted systematically by considering the program objectives of the AFCEE RNA demonstration, physiochemical properties of the CAH compounds, and other site-specific characteristics such as hydrogeology, land use assumptions, potential exposure pathways, and appropriate remediation goals. All of these factors will influence the technical effectiveness, implementation, and relative cost of technologies for remediating shallow groundwater underlying and migrating from the The remedial approaches and source removal technologies retained for development of remedial alternatives and comparative analysis include institutional controls, RNA combined with LTM, and soil vapor extraction (SVE) with potential aboveground treatment of extracted vapor. Groundwater extraction and treatment was not retained because it does not fulfill AFCEE's program objectives for waste minimization. However, the planned, limited groundwater extraction and treatment system to be installed during the spring/summer of 1998 by IT was considered in the implementation of remedial alternatives.

# INITIAL TECHNICAL IMPLEMENTABILITY SCREENING OF TECHNOLOGIES AND PROCESS OPTIONS FOR GROUNDWATER REMEDIATION TABLE 6.2

Retain	Yes	No	Yes	Yes		No	No		Yes	No		No				No		No
Relative Cost	Low	Low	Low	Low		Low	Moderate		Low	Moderate		Moderate				High		High
Effectiveness	Necessary for all remedia- tion strategies	Necessary	Necessary	Necessary		Not required at this site	Not required	at this site	Necessary	Low		Moderate				Low		Low
Implementability	New monitoring wells must be installed along the centerline of the plume lobes to properly characterize the progress of remediation. Sufficient space exists for additional wells.	The probable point-of-compliance likely has been impacted. The point-of-compliance (i.e., Long Branch Creek) will need to be monitored to confirm the progress of remediation.	As stated above, surface water of Long Branch Creek will be monitored.	The plume lies within the Base boundary, and land and groundwater use are under Base jurisdiction.		No production wells are known to exist in the current or predicted plume	No shallow groundwater is extracted from the plume area for any use.		Base public relations and environmental management offices have many information avenues through which to communicate to workers and residents.	Surface water has been impacted by discharge of contaminated	groundwater. Marsh terrain near creek may inhibit the proper installation of a trench. No likely human receptors downgradient.	No likely receptors downgradient from site and dissolved plume is	projected to recede. Long-term groundwater extraction does not comply with program objectives; however, short-term groundwater	extraction is currently proposed by the Base as an interim measure (IM)	within the source area until the viability of natural attenuation can be assessed.	Limited effectiveness. Contaminated groundwater would seek alternate	paths over, under, or around the walls enroute to the creek.	Limited effectiveness. Contaminated groundwater would seek alternate paths over, under, or around the walls enroute to the creek.
Process Option	Confirmation Wells	Sentry Wells	Surface Water	Land Use Control/Regulate	Well Permits	Seal/Abandon Existing Wells	Point-of-Use	Treatment	Meetings/ Newsletters	Interceptor	Trench Collection	Groundwater	Extraction			Slurry	Walls/Grout	Sheet Piling
Technology Type	Periodic Groundwater / Surface Water Monitoring	)		Groundwater Use Control					Public Education	Hydraulic	Controls	,				Physical	Controls	
General Response Action	Long-Term Monitoring			Institutional Controls						Containment of	Plume							

### TECHNOLOGIES AND PROCESS OPTIONS FOR GROUNDWATER REMEDIATION INITIAL TECHNICAL IMPLEMENTABILITY SCREENING OF TABLE 6.2 (continued)

	Ketaın	No		oN N	Yes	No No	SZ SZ	No No	No	No
	Kelative Cost	Moderate		Low	Low	Low	Moderate	High	Moderate	High (O&M)
:	Effective- ness	Moderate		Low	High	Low	Moderate	Moderate	High	Moderate
SHAW AFB, SOUTH CAROLINA	Implementability	Degradation of CAHs may be stimulated by allowing groundwater to flow	through a nutrient-rich zone or zero-valent iron. New, unproven technologies.	Nutrients and/or oxygen are injected within and/or downgradient of plume to limit plume migration by enhancing biodegradation and reducing CAH concentrations. Will inhibit anaerobic degradation of CAHs by oxygenating the groundwater. 1,1,1-TCA not aerobically biodegradable. Would be most effective downgradient of the source area. Not proven to be more effective than natural attenuation.	A combination of natural biological, chemical, and physical removal mechanisms which occur to varying degrees on every site. Groundwater sampling at OU-4 indicates that this is an ongoing remediation process.	Injection of air into contaminated aquifer creating a mass transfer of CAHs, specifically 1,1,1-TCA, into air bubbles and vadose zone. Similar to biosparging in effectiveness; however, more mass is transferred rather than destroyed. Most effectively used downgradient of the source so as not to upset anaerobic conditions promoting reductive dechlorination in the source area.	Source area of groundwater plume is pumped by installing submersible pumps in source area wells. Produces a large volume of water which requires additional treatment. However, will rapidly stop the migration of 1,1,1-TCA toward Long Branch Creek if sufficient pumping rates can be sustained (>20 gallons per minute) in the source area. Does not comply with program objectives.	High flow rates require excessive retention times and large reactors. CAHs are often volatilized in these systems.	Cost-effective technology for removing CAHs from groundwater at high flow rates. Potential permitting for air emissions.	Cost prohibitive for more concentrated CAHs. Creates a carbon disposal problem.
	Process Option	Biologically	Active Zones/Iron Filings Trench	Oxygen and/or Nutrient Enhanced Biodegradation (Biosparging)	Natural Attenuation	Air Sparging (Volatilization)	Vertical Pumping Wells	Bioreactors	Air Stripping	Activated Carbon
	Technology Type	Reactive/	Semi- Permeable Barriers	Biological	Chemical/ Physical		Groundwate r Extraction	Biological	Chemical/ Physical	
	General Response	Action In Situ	Groundwater			,	Source Removal/ Groundwater Remediation	Aboveground Groundwater Treatment		

## TECHNOLOGIES AND PROCESS OPTIONS FOR GROUNDWATER REMEDIATION INITIAL TECHNICAL IMPLEMENTABILITY SCREENING OF TABLE 6.2 (Continued)

General	Technology	Process Option	Implementability	Effectiveness	Relative	Retain
Response Action	Type				Cost	
Aboveground Groundwater Treatment		Direct Discharge to Industrial Waste Water Treatment Plant (IWWTP)	Implementable option only when an IWWTP is readily available and capable of handling contaminant and hydraulic loading. IWWTP not available for this site.	High	Low	No
		UV/Ozone Reactors	High flow rates require lengthy retention times and large, costly reactors.	Moderate	High	N <sub>o</sub>
Treated Groundwater Disposal	Discharge to IWWTP or Sanitary Sewer	IWWTP	Implementable option only when an IWWTP is available and capable of handling hydraulic loading. IWWTP is not available.	High	Low	No No
•		Sanitary Sewer	Implementable option only when access to a sanitary sewer exists and hydraulic loading is acceptable.	High	Low	No No
	Treated Groundwater	Vertical Injection Wells	Injection wells subject to clogging, high maintenance, and permitting.	Moderate	Moderate	%
	Reinjection	Injection Trenches	Requires large trenches and can be subject to injection well permitting.	Moderate	Moderate	No
	Discharge to Surface Waters	Storm Drains	Generally requires NPDES or other discharge permit.	High	Low	No
Source Removal/Soil Remediation	Excavation/ Treatment	Landfilling	Some excavation may economically be feasible if it can be shown that source area is generally less than 2,000 cubic yards. For reasons of risk and cost reduction, in-situ methods preferable when possible.	Moderate to High	Moderate	No No
		Biological Landfarming	Some excavation may be feasible at this site. For reasons of risk and cost reduction, <i>in-situ</i> methods preferable when possible.	Moderate to High	Moderate	No
		Thermal Desorption	Some excavation may be feasible at this site. For reasons of risk and cost reduction, <i>in-situ</i> methods preferable when possible.	Moderate to High	High	No
	In Situ	Bioventing	Air injection/extraction to increase soil oxygen levels and stimulate biodegradation of fuel residuals. Conductivity of site soils to air movement would be greater than to water movement. Might stimulate cometabolism of CAHs during the biodegradation of fuel contamination.	Moderate to High	Low	o V
		Soil Vapor Extraction	Air extraction to increase soil oxygen levels and extract VOCs from vadose zone. Conductivity of site soils to air movement would be greater than to water movement. May be subject to air emissions permitting.	High	Low to Moderate	Yes

## TECHNOLOGIES AND PROCESS OPTIONS FOR GROUNDWATER REMEDIATION INITIAL TECHNICAL IMPLEMENTABILITY SCREENING OF TABLE 6.2 (Concluded)

Retain	No	No	No	No	No
Relative Cost	High	High	Low to Moderate	High	Moderate to High
Effectiveness	Low	Moderate	Moderate	Moderate	Moderate to High
Implementability	Water and/or surfactant solution are forced through zones of residual contamination to enhance contaminant partitioning into the groundwater. Most effective in homogeneous sandy soils as present at OU-4.	Best suited for sites with >1 foot mobile LNAPL. Sandy soils at OU-4 will enhance effectiveness. Current LNAPL thicknesses observed to be < 1 foot.	Best suited for sites with >1 foot mobile LNAPL. Sandy soils at OU-4 will enhance effectiveness. Current LNAPL thicknesses observed to be < 1 foot.	Best suited for sites with thin saturated zones where excessive groundwater will not be pumped.	Best suited for low-permeability/low-transmissivity sites. Residual and mobile LNAPL, soil vapor, and groundwater can be extracted. Bioslurping is a subset of this technology.
Process Option	Soil Washing	Dual-Pump Systems	Skimmer Pumps, Bailers, Wicks	Total Fluids Pumping	Vacuum-enhanced extraction
Technology Type	In-Situ (continued)	Mobile LNAPL Recovery			
General Response Action	Source Removal/Soil Remediation (continued)				

Plume capture downgradient from the source area and prior to discharge to Long Branch Creek also could be accomplished by installing an air sparging curtain or iron filings trench (permeable reaction wall). An iron filings trench was installed below the upper terrace near Long Branch Creek in November 1998 to mitigate discharge of dissolved contaminants to the creek (see Appendix E for location). With an iron filings trench, groundwater contaminated with CAHs flows through granular, zero-valent iron filings placed in a trench, which is excavated perpendicular to the axis of groundwater flow. Dissolved CAHs are degraded through a series of less-chlorinated intermediates to non-toxic, non-chlorinated end products. However, iron filings trenches are a new, relatively unproven technology. The cost for the iron filings backfill can range from \$400 to \$450 per ton (Focht et al., 1996) (approximately \$740 to \$840 per cubic yard), and the effective lifetime of an iron filings trench is not known. There is a potential for precipitation of dissolved minerals within the trench due to pH changes across the trench boundary, resulting in clogging. Therefore, an iron filings trench may require periodic rehabilitation or replacement. Due to the unproven nature of this technology, the potential for high long-term maintenance costs, the lack of source treatment to reduce plume longevity, and the potential for surface discharge of groundwater, an iron filings trench was not selected in any alternative. Nonetheless, the current permeable reaction wall at OU-4 is included in Alternative 1, and should be evaluated as data regarding the effective lifetime and long-term cost of this technology are assessed.

An air sparging curtain could be constructed by installing one or more horizontal, perforated pipes in a gravel-filled trench or numerous vertical wells in a row perpendicular to plume migration. Air injected into the pipes or wells rises through the contaminated groundwater and strips volatiles from the groundwater, transferring them into the vadose zone. If necessary to meet air quality standards, soil vapor extraction (SVE) and aboveground treatment systems could be installed to capture and treat the stripped volatiles. Some researchers have cast doubt on the long-term effectiveness of air sparging. Problems such as channeling, which consists of preferential migration of injected air along specific (more permeable) flow paths rather than uniform air dispersal in the zone surrounding the perforated pipe, have been cited. The occurrence of channeling could reduce the integrity of the sparging curtain in some areas, and allow contaminants to migrate past the curtain. However, the potential for channeling in a homogenous, gravel-filled trench is reduced. Air sparging in the relatively homogeneous sandy soils below the upland terrace near the toe of the CAH plume should be effective without the aid of a gravel trench.

A small percentage of the injected air in an air sparging system would most likely become dissolved in the groundwater, increasing the DO content within the sparging trench and for some distance downgradient from the system. As described in Section 4.4.6.1, DO represents an alternate electron acceptor that, if used preferentially over CAHs, may inhibit the occurrence of reductive dechlorination within the northeast or southeast/east plume lobes. However, less-chlorinated biodegradation daughter products from anaerobic and abiotic processes (e.g., DCE isomers, VC, DCAs, and CB) would potentially be aerobically biodegraded in part to the increased oxygen delivery to the groundwater aquifer. 1,1,1-TCA is not known to be aerobically biodegradable. Air sparging is potentially feasible at the OU-4 site, but was not retained for detailed evaluation due to planned groundwater extraction and treatment near monitoring wells MW-112 and MW-117 by IT Corporation. Air sparging or an iron fileings trench could be reconsidered at a later date if groundwater pump and treat

operations do not reduce/control groundwater contamination at the site to the satisfaction of the SCDHEC.

Contaminated soils near the plume hotspot at MPC could be excavated for the most rapid reduction of the contaminant plume. Modeling results indicate that groundwater velocities are sufficiently high that a rapid decrease in the leachable mass of CAHs in the source area would result in dramatic reduction of CAH concentrations across the plume within 2 to 3 years. However, the extent of soil contamination would first need to be further defined with a soil gas survey and/or additional soil sampling. Unless the volume of contaminated soils in the source area was less than 2,000 cubic feet, the benefits of soil excavation may be offset by the costs associated with contaminated soil In addition, soil excavation conflicts with excavation, treatment, and disposal. AFCEE's waste minimization requirements. The mobile LNAPL detected at the site and much of the residual soil contamination appears to border the water table at a depth of approximately 12 feet bgs, and potentially extends over several thousand square feet. Therefore, the potentially excessive cost of soil excavation combined with the potentially large volume of soil that would require ex situ treatment and disposal, precluded further evaluation of this option in this TS.

### 6.3 BRIEF DESCRIPTION OF REMEDIAL ALTERNATIVES

This section describes how remedial technologies retained from the screening process were combined into two remedial alternatives for OU-4. Sufficient information on each remedial alternative is provided to facilitate a comparative analysis of effectiveness, implementability, and cost in Section 6.4.

### 6.3.1 Alternative 1--RNA and Institutional Controls with Long-Term Groundwater and Surface Water Monitoring

RNA is achieved when naturally occurring attenuation mechanisms bring about a reduction in the total mass of a contaminant in the soil or dissolved in groundwater. RNA results from the integration of several subsurface attenuation mechanisms that are classified as either destructive or nondestructive. Destructive attenuation mechanisms include biodegradation, abiotic oxidation, and hydrolysis. Nondestructive attenuation mechanisms include sorption, dilution (caused by dispersion and infiltration), and volatilization. In some cases, RNA will reduce dissolved contaminant concentrations below numerical concentration goals intended to be protective of human health and the environment. As indicated by the evidence of RNA described in Section 4, these processes are occurring at OU-4.

Implementation of Alternative 1 would require the use of institutional controls such as land use restrictions and LTM. Land use restrictions may include placing long-term restrictions on soil excavation in the source area, on groundwater well installations within and downgradient from the plume area, and on use of Long Branch Creek east of the site. The intent of these restrictions would be to reduce potential receptor exposure to contaminants by legally restricting activities within areas affected by site-related contamination.

Public education on the selected alternative would be developed to inform Base personnel and residents of the scientific principles underlying RNA. This education

could be accomplished through public meetings, presentations, press releases, and posting of signs where appropriate. Periodic site reviews also could be conducted using data collected during the LTM program. The purpose of these reviews would be to evaluate the extent of contamination, assess contaminant migration and attenuation through time, document the effectiveness of institutional controls at the site, and reevaluate the need for additional remedial actions at the site.

The numerical model described in Section 5 was used to simulate the long-term effectiveness of this alternative. The simulation setup and results are described in detail in Section 6.4.1.

### 6.3.2 Alternative 2—SVE of the 1,1,1-TCA Source, RNA, and Institutional Controls with Long-Term Groundwater and Surface Water Monitoring

Alternative 2 is identical to Alternative 1 except that a SVE system would be installed within the main source area creating the east/southeast 1,1,1-TCA contaminant plume lobe (near MPC). This system would treat the residual LNAPL source at this location and also promote volatilization of VOCs from mobile LNAPL, thereby treating the groundwater "hot spot" near MPC and reducing discharge to Long Branch Creek.

According to Cohen and Mercer (1993), dissolved contaminant concentrations greater than 1 percent of the aqueous solubility are highly suggestive of NAPL presence. For 1,1,1-TCA, 1 percent of the aqueous solubility is 13,600  $\mu$ g/L. Therefore, the presence of dense non-aqueous phase liquid (DNAPL) cannot be ruled out.

Despite the desirability of source reduction/removal, it may be technically difficult to achieve given the fact that DNAPL may be present beneath OU-4 and that significant LNAPL contamination may reside in a thin layer bordering the water table. However, simulation of this alternative using the numerical MODFLOW/MT3D<sup>96®</sup> model indicates how the plume would be affected in the future if the source could be at least partially remediated. Without source removal/reduction, implementation of other remedial actions (e.g., proposed groundwater pump-and-treat as an interim remedial action) would be complicated by the fact that the remedial system(s) would have to operate indefinitely due to the potential longevity of the source.

SVE is applicable for OU-4 because soils are highly permeable, which would allow efficient air movement and oxygen transport. Moderate seasonal changes in water table elevation at the site (historically observed to fluctuate by as much as two feet) would expose more contaminated soils to the effects of SVE during low-water periods and cause an increase in vapor flow to the extraction well. The induced vapor flow volatilizes adsorbed and free-phase contaminants and removes them from the liquid phase.

Extracted soil vapor could be released to the atmosphere if contaminant concentrations are below state discharge requirements. Alternatively, vapor could initially be treated through catalytic oxidation. The catalytic oxidation system would be used to treat initially high VOC concentrations at the site. After VOC concentrations were reduced, the catalytic oxidation system could be taken off-line and replaced by granular activated carbon (GAC) beds. The catalytic oxidation system purifies

contaminated air streams by heating and then destroying the contaminant vapors by contact with an oxidation catalyst. The SVE blower would discharge VOC laden soil gas through a heat exchanger where it is preheated prior to transfer to the catalyst chamber. Once heated, the vapor laden air mixture passes through the catalyst for exothermic destruction. The gases are cooled and scrubbed for subsequent treatment of hydrochloric acid (HCl), a byproduct of catalytic destruction of chlorinated VOCs.

After several months of SVE operation with a catalytic oxidizer, soil gas concentrations may be low enough to use GAC beds for emission treatment. GAC removes VOCs from soil gas by sorbing the VOC to the surface area of the GAC. GAC treatment is more economical for low VOC concentration waste streams and can achieve good removal efficiencies. Periodic replacement of the GAC beds would be required as their available surface area becomes saturated.

A small pilot-scale study would need to be implemented to determine the appropriate number of extraction wells, well radius, available screen area, vacuum configuration, and extraction rates of an SVE system. In order to evaluate the effectiveness of this remedial option, the numerical model was used to simulate the reasonable effects of source reduction through SVE in the source area of the east/southeast source area of the CAH plume. The simulation setup and results are described in detail in Section 6.4.2.

### 6.4 EVALUATION OF ALTERNATIVES

This section provides a comparative analysis of each of the remedial alternatives based on the effectiveness, implementability, and cost criteria. A summary of this evaluation is presented in Section 6.5.

### 6.4.1 Alternative 1--RNA and Institutional Controls with Long-Term Groundwater and Surface Water Monitoring

### 6.4.1.1 Effectiveness

The Visual MODFLOW and MT3D<sup>96®</sup> codes were used to simulate plume migration if no engineered remedial actions are performed. Model OU4MODA was run for 100 years beyond 1997. Although it is realistic to assume that the source strength will decrease over time due to the effects of leaching and weathering/degradation, the rate of decrease cannot be predicted with confidence due to the lack of information regarding the nature and location of the source. Cohen and Mercer (1993) state that factors influencing NAPL dissolution and eventual depletion include the effective aqueous solubility of NAPL components, groundwater velocity, NAPL-water contact area, and the molecular diffusivity of the NAPL chemicals in water. The actual dissolution of NAPL will generally slow with time due to aging and reduction of the NAPL-water contact area (Powers et al., 1991).

The presence of LNAPL (and potentially DNAPL) at OU-4 could conceivably result in significantly elevated dissolved concentrations of chlorinated solvent, specifically 1,1,1-TCA, for generations. As mentioned in Section 4.1, a cosolvenated fuel/solvent LNAPL was identified at monitoring point MPC, whereas a fuel LNAPL was identified at monitoring point MPB. A product sheen of unknown type was observed at monitoring point MPD. Each of these LNAPL locations is susceptible to weathering,

especially considering the warm surface temperatures, the close proximity of the groundwater surface to the ground surface, and high soil permeabilities which enhance volatilization and precipitation-induced leaching.

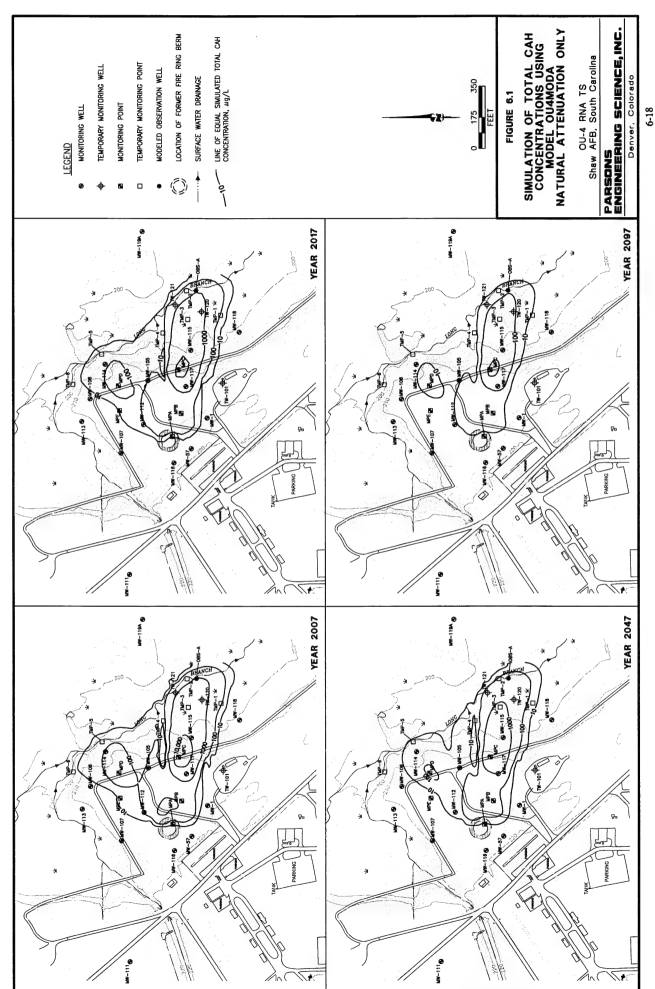
For prediction purposes, the contaminant sources at OU-4 were predicted to decrease at a geometric rate of 3 percent per year starting in May 1997. At this rate, the source strengths decreased by 26 percent by simulation year 10 (calendar year 2007) and by 46 percent by simulation year 20 (calendar year 2017). This source reduction scenario reflects both the potential for significant dissolved groundwater contamination for many years, and the likelihood that the source strength will decrease over time.

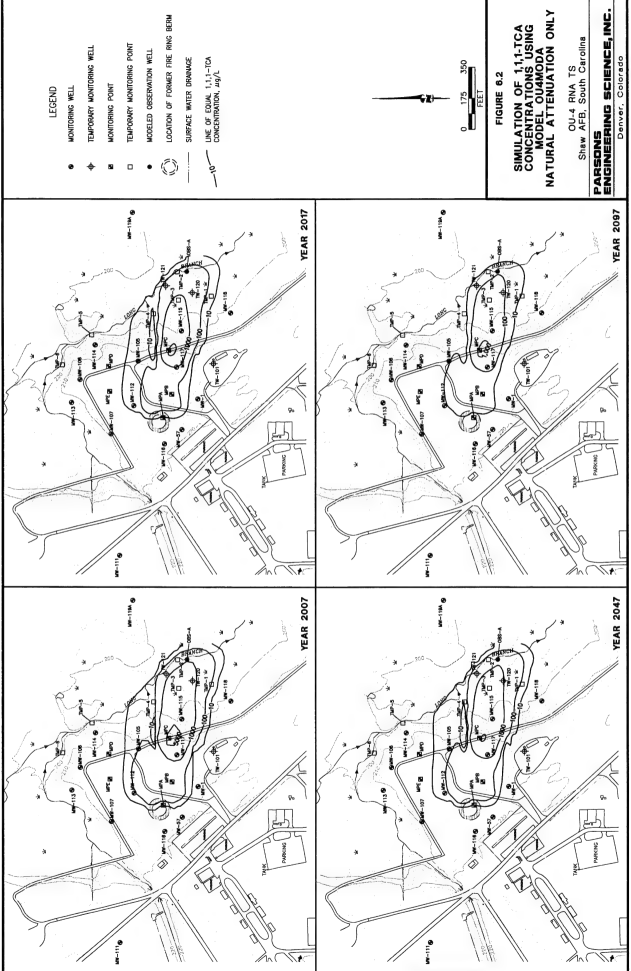
Figure 6.1 presents the migration of the calibrated total CAH plume over time predicted by Model OU4MODA under the influence of RNA alone. Rapid groundwater velocities cause the plume to discharge to Long Branch Creek before the CAHs can be completely biodegraded. As mentioned in Section 4, the approximate migration time for a contaminant molecule to travel from the former fire training berm to the creek estimated to be approximately 2 years. This migration rate results in rapid stabilization of the plume and little variation in the plume length or width is predicted during much of the 100-year simulation period. Observed CAH concentrations within the source area and downgradient at the edge of Long Branch Creek in May 1997 are predicted to be maximum groundwater concentrations. The model predicts that these concentrations will diminish (at monitoring point MPC) to 26,040  $\mu$ g/L, 19,190  $\mu$ g/L, 7,700  $\mu$ g/L, and 1,440  $\mu$ g/L for calendar years 2007, 2017, 2047, and 2097, respectively. Similarly, maximum dissolved CAH concentrations at Long Branch Creek are predicted to decrease steadily to 101  $\mu$ g/L by calendar year 2097.

Figure 6.2 depicts the simulated decreases in 1,1,1-TCA concentrations over the same 100-year time period. 1,1,1-TCA was assumed to represent approximately 56 percent of the total CAH plume mass at OU-4. This ratio was determined by calculating the weighted average of 1,1,1-TCA relative to total CAH contamination for each well where 1,1,1-TCA was detected. The total CAH plume was then converted to 1,1,1-TCA concentrations using the 56 percent ratio. Calculations are included in Appendix C. By using a weighted average, the high 1,1,1-TCA concentrations in the source area cause the simulated 1,1,1-TCA concentrations at the periphery of the plume to be biased high. The model indicates that maximum 1,1,1-TCA concentrations in groundwater will not achieve the current interim remedial goal of 1,000  $\mu$ g/L at the site within the next 95 years at a geometric source weathering rate of 3 percent per year. Figure 6.3 is a graphic showing predicted 1,1,1-TCA groundwater concentrations versus time in the source area (MPC) and near Long Branch Creek (OBS-A).

### **6.4.1.2** Technical and Administrative Implementability

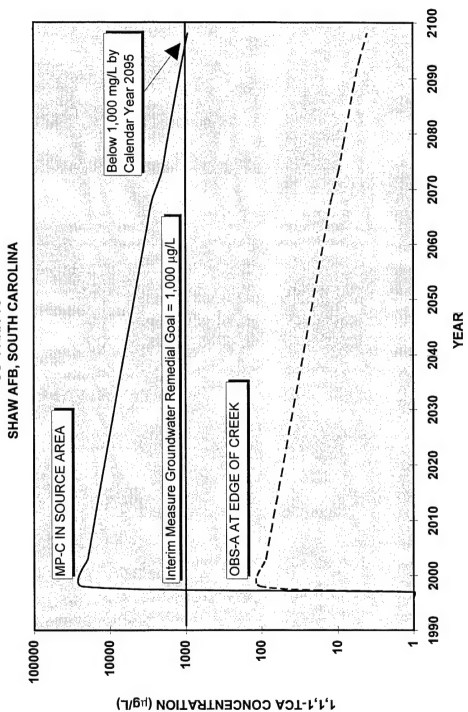
Alternative 1 is not technically difficult to implement. Installation of LTM and POC wells and periodic groundwater and surface water sampling and analysis are standard procedures. Long-term management efforts would be required to ensure that proper sampling procedures are followed. Periodic site reviews should be conducted to confirm the adequacy and completeness of LTM data and to verify the effectiveness of





2:/E2/cad/AFCEE/722450/SHAW/98dn0123.dwg, 04/30/98 at 08:25

FIGURE 6.3
PLOT OF 1,1,1-TCA CONCENTRATIONS VERSUS TIME
FOR MODEL OU4MODA
OU4 RNA TS



this remediation approach. However, there may be administrative concerns associated with long-term enforcement of groundwater use restrictions. Given the assumptions regarding the future magnitude and duration of the CAH source, specifically 1,1,1-TCA, that are incorporated into the predictive models, substantially elevated 1,1,1-TCA concentrations are predicted to persist in groundwater for nearly a century. Future land use within the plume area may be impacted by leaving contaminated soil and groundwater in place. Regulators and the public will have to be informed of the benefits and limitations of the RNA option, but educational programs are not difficult to implement.

### 6.4.1.3 Cost

The present-worth cost of implementing Alternative 1 for a 20-year period beginning in 1998 is summarized in Table 6.3. Included in the total present-worth cost of \$232,110 are the estimated costs for installing additional LTM and sentry wells, performing the recommended groundwater and surface water monitoring (see Section 7), maintaining institutional controls, public education, project management, and reporting. Cost calculations are contained in Appendix E.

### 6.4.2 Alternative 2 - SVE of the 1,1,1-TCA Source, RNA, and Institutional Controls with Long-Term Groundwater and Surface Water Monitoring

### 6.4.2.1 Effectiveness

Model OU4MODA, used to simulate the effectiveness of Alternative 1 (Section 6.4.1.1), was modified to include the effects of a SVE system operating in the source area of the east/southeast CAH plume lobe near MPC. The SVE system was assumed to begin operation in the fall of 1998 after proposed pilot scale testing in the summer of 1998; the system was assumed to operate for a total of 5 years until calendar year 2003.

The resulting model, termed OU4MODB, assumes annual geometric source reduction of 8-percent during years 1 and 2 of SVE system operation, 5-percent reduction during years 3 through 5, and 3-percent reductions for the remainder of the model simulation. At this rate, 1,1,1-TCA source area concentrations are predicted to undergo reductions of 38 percent by 5 years and 54 percent by 20 years. The model assumes that SVE will not be implemented at other potential source areas in the former fire training berm and northeast of the berm near monitoring point MPD. These source areas were assumed to naturally weather at a geometric rate of three percent per year.

It should be noted that the effectiveness of a SVE system at remediating the CAH source cannot be predicted with accuracy because the nature and extent of the source is unknown. If the source of the east/southeast plume lobe primarily borders the water table, SVE would be most effective during seasonal groundwater lows. Groundwater elevations measured in May 1997 were approximately 2 feet lower than groundwater elevations measured in March 1993. This suggests that the NAPL layer may have been thicker in May 1997 because more residual contamination was exposed that could result in NAPL formation. However, temporal water table fluctuations are beneficial to SVE because more residual contamination can be exposed to SVE than if the water table elevation did not fluctuate. Therefore, the results of the model simulations incorporating SVE are speculative and the effectiveness of SVE could be greater or less

### TABLE 6.3 ESTIMATED ALTERNATIVE 1 COSTS OU4 RNA TS SHAW AFB, SOUTH CAROLINA

Capital Costs	Present Worth Cost
Design/construct three LTM wells and one sentry well in 199	\$12,450
Monitoring Costs	Present Worth Cost
Conduct annual groundwater monitoring of 11 wells and 6 surface water stations from 1998 to 2002	\$97,264
Conduct biennial groundwater monitoring of 11 wells and 6 surface water stations from 2003 to 2018	\$57,390
Site management (maintain institutional controls/public education) (20 years)	\$65,010
Total Present Worth of Alternative 1 a/	\$232,110

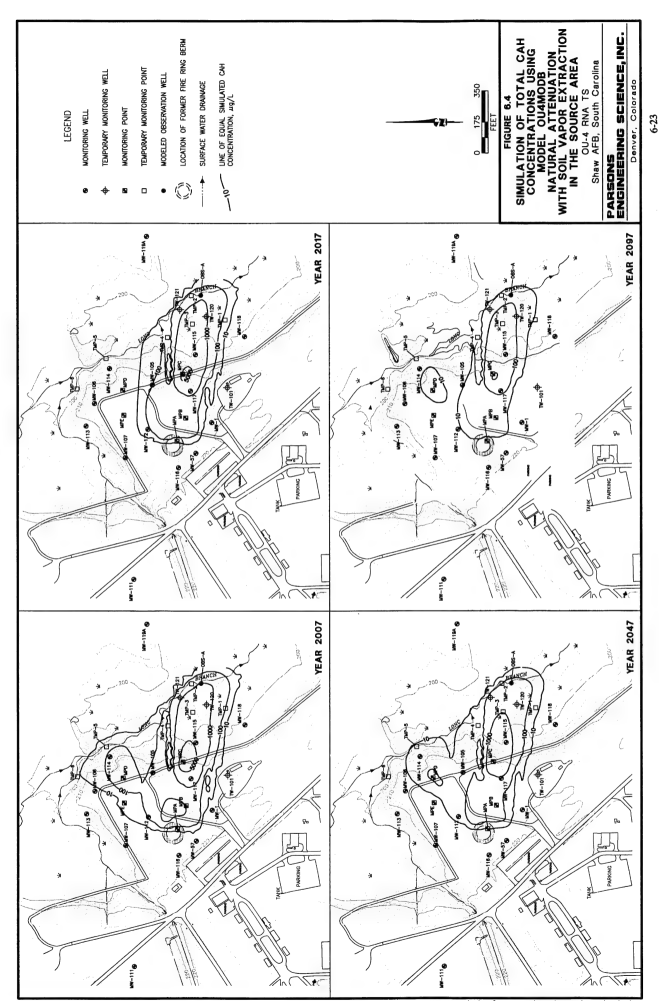
a/ Based on an annual adjustment factor of 7 percent (USEPA, 1993).

Note: Costs assume that well installation and LTM are performed by local (Sumter/Columbia) personnel.

than predicted. However, the simulations indicate how engineered source reduction, if feasible, would affect the CAH and 1,1,1-TCA concentrations.

The results of a SVE pilot test performed in the former fire training berm in February 1993 indicated that significant soil vacuums were generated at the site with a pore volume turnover rate of at least 24 per day (Rust E&I, 1993). The radius of influence of vertical SVE extraction wells was limited due to the presence of a perched groundwater layer and confining clay sequence at approximately three to five feet bgs. The results of this study suggested that SVE is possible at the site, but its effectiveness may be limited by the geology beneath the fire training berm area. Because sandy soils are prevalent near MPC, an SVE system installed at this location would be expected to achieve better NAPL removal rates than measured at the visible fire training berm.

The CAH plume migration simulated by Model OU4MODB is shown on Figure 6.4. SVE reduces the maximum dissolved CAH concentration in the plume from 30,240  $\mu$ g/L in 1997 to 13,530  $\mu$ g/L by year 2018; therefore, maximum CAH concentrations within the plume are reduced more rapidly than in the Alternative 1 simulations (Section 6.4.1). Plume concentrations and extents tend to be similar to those predicted in model OU4MODA, except that overall concentrations are reduced approximately 12 percent faster by calendar year 2008 in model OU4MODB.



Simulated 1,1,1-TCA concentrations derived from model OU4MODB are shown on Figure 6.5. Although 1,1,1-TCA concentration reductions are accelerated in this prediction relative to Model OU4MODA, the areal extent of the plume through time is very similar to that predicted by Model OU4MODA (Figure 6.2). Similar to CAH concentrations, 1,1,1-TCA concentrations diminish approximately 12 percent faster than in Alternative 1 by calendar year 2008 (5 years of SVE and 5 years of normal weathering). 1,1,1-TCA concentrations are not predicted to decrease below 1,000  $\mu$ g/L at the site within 90 years. Figure 6.6 is a graphic showing predicted 1,1,1-TCA groundwater concentrations versus time in the source area (MPC) and near Long Branch Creek (OBS-A).

### 6.4.2.2 Technical and Administrative Implementability

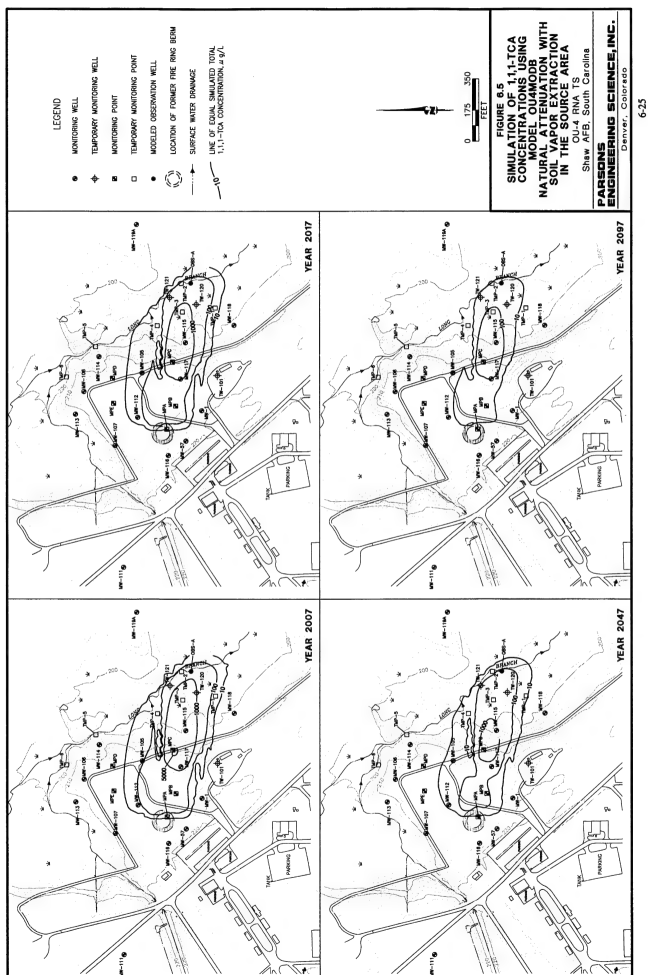
The implementability issues described for Alternative 1 in Section 6.4.1.2 also are applicable for Alternative 2. However, administrative concerns regarding the length of LTM and institutional controls may be lessened due to more rapid depletion of significantly elevated dissolved TCE concentrations. Regulatory and community acceptance of this alternative should be more positive than with Alternative 1 due to the more aggressive plume remediation activities.

Alternative 2, as simulated by the numerical model, should not be difficult to implement technically or administratively. However, the nature and extent of the 1,1,1-TCA source near MPC, specifically the LNAPL source, has not been fully defined. Additional site investigation would need to be performed to properly size and locate a SVE system.

The SVE system simulated by model OU4MODB would require construction of approximately 5 SVE wells (assuming a radius of influence of 50 feet for each well) in the vicinity of monitoring point MPC, and installation of an aboveground vacuum blower, catalytic oxidation and GAC emission treatment systems, and associated piping. SVE is assumed to be limited to the area adjacent to monitoring point MPC where fuel/solvent LNAPL was detected.

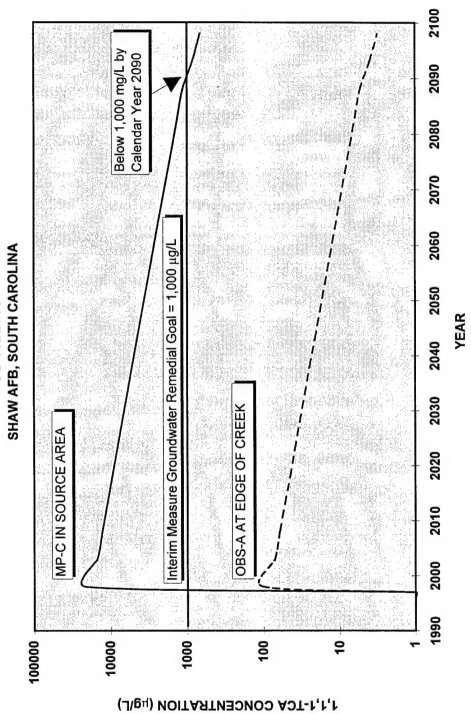
The requirement for vapor treatment systems (e.g., catalytic oxidation/GAC systems) potentially may be omitted from the design process if appropriate land-use controls can be maintained and vapors may be emitted to the atmosphere. Otherwise, it is recommended that a catalytic oxidation unit with an air scrubber for HCl initially be used at the site to reduce emissions until influent concentrations to the SVE blower were low enough to utilize a cheaper GAC treatment system. Catalytic oxidation and GAC systems are generally reliable when properly maintained. Standard mechanical and electrical construction and equipment would be used; special construction techniques and special-order equipment should not be necessary. All equipment needed for this alternative is commercially available.

Assuming a 5-year-long operating period, the above ground vacuum blower(s) may have to be replaced once over the life of the SVE system. The most significant maintenance requirement for this system will be regular maintenance of the catalytic oxidation system during the (estimated) first 3 months of initial SVE system operation and of the GAC system during the following years to ensure that air emissions standards are attained. Blower systems are very reliable and require only minimal



05/12/98 at 11:03

FIGURE 6.6
PLOT OF 1,1,1-TCA CONCENTRATIONS VERSUS TIME
FOR MODEL OU4MODB
OU4 RNA TS



maintenance. Air filters provide blower protection and generally require replacement every 90 to 180 days. Installation and operation of a full-scale SVE system would require an increased commitment of man-hours and other resources to maintain and monitor the system. Weekly system checks are recommended, and operating data such as vacuum pressure and flow rate should be recorded. Periodic vapor samples would be required to demonstrate compliance with emissions requirements.

Performance of an SVE pilot test would be desirable to obtain design parameters to facilitate design of a full-scale system. The pilot test could potentially be performed using new monitoring wells installed for groundwater pump and treat by IT (SCDHEC, 1998).

Administrative implementation of this alternative will require that appropriate air discharge permits are obtained, and that future development plans protect the SVE system. All system components (above- and below-grade) should be protected against damage in the event that construction or maintenance work is performed in the area.

### 6.4.2.3 Cost

The estimated capital and operating costs of Alternative 2 are shown in Table 6.4. The total present worth cost of Alternative 2 is approximately \$593,590. The cost of Alternative 2 is greater than Alternative 1 due to the addition of the SVE system. It is assumed that the SVE system would be operated for 5 years after installation. Annual LTM would continue for a total of 5 years to build a historical groundwater and surface water quality database, and biennial LTM would be performed for at least an additional 15 years to monitor temporal changes in dissolved CAH concentrations. Cost calculations are contained in Appendix E.

### 6.5 GROUNDWATER PUMP-AND-TREAT AS AN INTERIM MEASURE TO MONITORED NATURAL ATTENUATION

Shaw AFB currently is seeking monitored natural attenuation as a long-term remedy at OU-4, to be supported, in part, by this TS. However, the SCDHEC has mandated that an IM be implemented to mitigate groundwater migration until monitored natural attenuation can be evaluated at the site. The currently proposed IM for site OU-4 is a groundwater pump-and-treat system to be installed by IT Corporation during the spring/summer of 1998. It is assumed for purposes of this report that the planned groundwater extraction and treatment will be conducted for several years until sufficient groundwater and surface water data can be produced supporting natural attenuation and stabilization/recession of the groundwater plume.

The effects of groundwater pump-and-treat within the source area near monitoring point MPC in combination with natural attenuation were evaluated in model OU4MODC. Specific design information regarding the implementation of the groundwater extraction and treatment system, including anticipated pumping rates, well construction details, and precise locations, were not available at the time of this report, but were being prepared (IT, 1998). Therefore, groundwater pump and treat in model OU4MODC was simulated by a single well located adjacent to MPC and screened from several feet above the water table to approximately 20 feet below the water table.

### TABLE 6.4 ESTIMATED ALTERNATIVE 2 COSTS OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

Capital Costs	<b>Present Worth Cost</b>
Design/construct three LTM wells and one sentry well in 199	8 \$12,450
Design/install SVE system in 1998	\$115,860
Monitoring and O&M Costs	Present Worth Cost
Conduct annual groundwater monitoring of 11 wells and 6 surface water stations from 1998 to 2002	\$97,260
Conduct biennial groundwater monitoring of 11 wells and 6 surface water stations from 2003 to 2018	\$57,390
Site management (maintain institutional controls/public education) (20 years)	\$65,010
Operate/maintain SVE system (5 years)	\$245,620
Total Present Worth of Alternative 2 a/	\$593,590

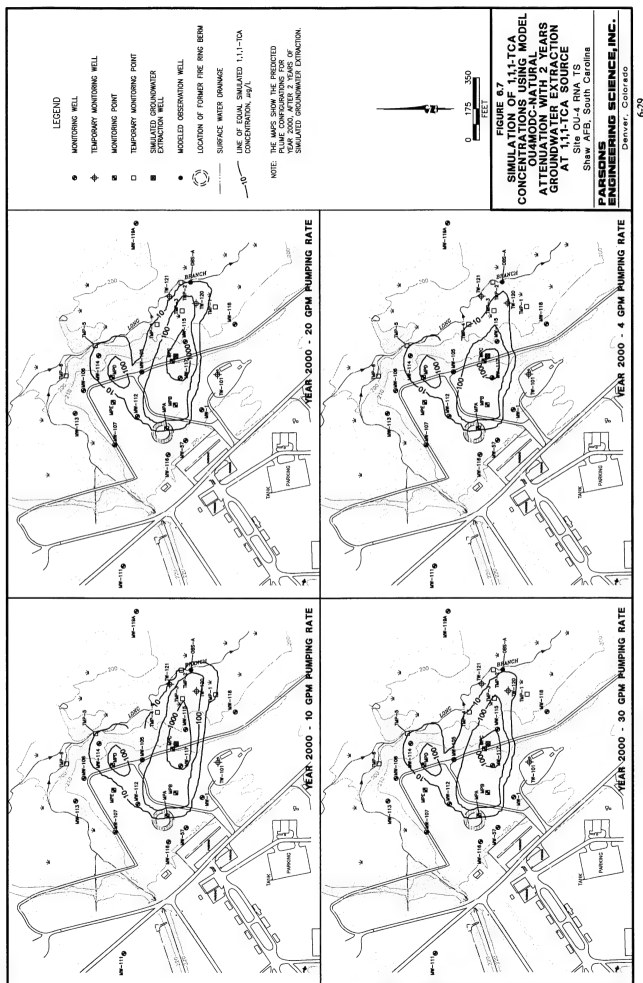
a/ Based on an annual adjustment factor of 7 percent (USEPA, 1993).

Note: Costs assume that well installation, LTM, and O&M are performed by local (Sumpter/Columbia) personnel.

Simulated pumping rates ranging from 10 to 40 gpm were assumed to be feasible given the sandy, relatively permeable soils underlying the site.

Figure 6.7 presents results from model OU4MODC for the year 2000 at selected pumping rates from 10 to 40 gpm. At the lower pumping rate of 10 gpm, the simulated groundwater plume is similar to the 1997 plume (Figure 4.7), and concentrations of 1,1,1-TCA approaching or exceeding 100  $\mu$ g/L at the edge of Long Branch Creek may be possible (the maximum 1,1,1-TCA concentration measured adjacent to the creek in May 1997 was 78.6  $\mu$ g/L at TMP-2). At pumping rates of 20 gpm and above, the 1,1,1-TCA plume begins to recede and downgradient migration of the plume is inhibited by the pumping system. The simulated pumping rate of 40 gpm in model OU4MODC prevented 1,1,1-TCA concentrations greater than 1,000  $\mu$ g/L from migrating farther than 75 feet downgradient from the pumping well.

The model simulations indicate that beneficial results could be obtained by the implementation of a pump-and-treat system. However, the use of this type of system is costly and requires a high degree of operation and maintenance. Furthermore, the use of groundwater pump and treat will alter the hydraulics (and possibly also the geochemistry) of the groundwater system in the plume area, and current groundwater



04/AFCEE\722450/SHAW\98dn0155.dwg, 05/12/98 at 10:43

velocity estimates and estimated biodegradation rates used in the predictive models OU4MODA and OU4MODB may be invalidated. The groundwater pump and treat system also may have the negative effect of transporting oxygenated water from outside the source area to within the source area near the extraction well. The addition of oxygenated water to the source area would potentially suppress the microbial communities responsible for reductive dechlorination of CAHs or alter the redox conditions necessary for abiotic dehydrohalogenation of 1,1,1-TCA to 1,1-DCE. It is therefore recommended that groundwater extraction and treatment be used as only a temporary IM (as planned), replaced by remedial alternatives 1 or 2 as discussed in Section 6.6.

### 6.6 RECOMMENDED REMEDIAL APPROACH

Two alternatives have been evaluated for remediation of the shallow groundwater at OU-4. Components of the alternatives evaluated include SVE, aboveground treatment of extracted vapors, RNA with LTM of groundwater and surface water, and institutional controls. The alternatives were developed to maximize the benefit of RNA in the overall remedial process and to supplement the remedial strategy (if needed) with proven low-cost, non-intrusive source treatment technologies to shorten LTM efforts. Therefore, implementation of the reactive barrier walls installed at the site in 1998 was not evaluated because they remain an unproven technology. The reactive barrier walls will help to limit downgradient contaminant migration, but will not treat the contaminant source and, therefore will not reduce the length of an OU-4 LTM program. Table 6.5 summarizes the results of the evaluation based upon effectiveness, implementability, and cost criteria.

Alternative 1 makes maximum use of natural attenuation mechanisms to reduce plume migration and toxicity. Alternative 2 would reduce plume longevity and provide additional protection against further plume migration, but would still rely on RNA to reduce plume toxicity in the downgradient portions of the affected area. Implementation of Alternative 2 would potentially decrease the time frame required for remediation, but would require greater capital and O&M expenditures.

Each of the alternatives has implementability concerns. Implementation of Alternative 1 may require a very lengthy LTM period, and maintenance of institutional controls in the plume area would be required. However, if future LTM results indicate that the plume is stable or receding, and that surface water standards will not be exceeded, then the frequency of LTM sampling events could be reduced. installation of reactive barrier walls downgradient of the source is expected to help reduce the frequency of LTM sampling. SVE of the source area could further reduce the duration of LTM, but the effectiveness of SVE may be limited if most of the source is comprised of a thin NAPL layer, which may be seasonally water-saturated. SVE may potentially require more than 5 years to remediate the source if these conditions exist. In addition, if DNAPL is present, it would be unaffected by the SVE system and would continue to be a long-term source of dissolved groundwater contamination. However, without concurrent source reduction, any LTM, planned groundwater extraction and treatment, or maintenance of a reactive barrier wall would have to be performed indefinitely. Implementation of an SVE system (Alternative 2) instead of Alternative 1 (no source treatment) may be more acceptable to the public and regulatory

## SUMMARY OF REMEDIAL ALTERNATIVES EVALUATION TABLE 6.5

## OU-4 RNA TS SHAW AFB. SOUTH CAROLINA

	SHAW AFB, SOUTH CAROLINA	H CAROLINA	
Remedial Alternative	Effectiveness	Implementability	Present Worth Cost Estimate
Alternative 1 - Natural Attenuation - Long-Term Monitoring - Institutional Controls	Contaminant mass, volume, and toxicity will be slowly reduced over time, but due to the potentially large LNAPL source, the dissolved plume may persist for 95 years, necessitating a potentially lengthy period of LTM and institutional controls.	Readily implementable. Groundwater water quality monitoring required for an estimated minimum of 20 years, and potentially much longer. Institutional controls, including land and groundwater use controls, may be required for 95 years due to the persistence of elevated dissolved 1,1,1-TCA concentrations. Minimal exposure to potential receptors if institutional controls are implemented. If significant plume migration occurs and completion of exposure pathways is possible, then additional remedial work may be required.	\$232,110
Alternative 2 - Soil Vapor Extraction - Natural Attenuation - Long-Term Monitoring - Institutional Controls	Similar to Alternative 1, with the addition of a Soil Vapor Extraction (SVE) system to remove mobile and residual LNAPL. Contaminant mass, volume, and toxicity will be reduced more rapidly than in Alternative 1. However, even if the source is substantially reduced, significantly elevated dissolved CAH, specifically 1,1,1-TCA, concentrations may still persist for 90 years, and downgradient plume migration may continue to reach Long Branch Creek, depending on the effectiveness of SVE and actual biodegradation rates.	Installation/operation of the SVE system simulated by the numerical model should not pose significant implementability concerns. May require the use of catalytic oxidation followed by granular activated carbon treatment to reduce emissions. Long-term site management, groundwater use controls, and monitoring would be required as elevated 1,1,1-TCA concentrations may persist for 90 years. If significant plume migration occurs and completion of exposure pathways is possible, then additional remedial work may be required.	\$593,590

6-31

agencies because it represents the more aggressive remedial approach intended to reduce the duration of LTM.

The estimated present worth cost for Alternative 2 is substantially higher than for Alternative 1 due to the addition of engineered remedial actions. Operation and maintenance of an SVE system would require a more significant annual expenditure of funds during the operational lifetime of this system.

On the basis of this evaluation, Alternative 1 is selected with continued operation of the installed reactive barrier walls. The reactive barrier walls were installed as part of a separate effort, but their partial capture of the CAH plume will affect the sampling frequency of the LTM program because less contamination will reach Long Branch Available data indicate that the CAH and 1,1,1-TCA plumes are either stabilized or diminishing slowly due to RNA alone, and that 1,1,1-TCA concentrations decrease by more than 2 orders of magnitude between the source area and Long Branch Data is unavailable to describe current CAH reductions, but the combined affect of RNA and the barrier walls are expected to mitigate contaminant migration into Long Branch Creek. However, the degree to which RNA and the reactive barrier walls will continue to prevent significant downgradient migration of the plume is not known with certainty and will become more apparent as successive LTM sampling events are performed. A reduction of the sampling frequency for LTM might be reached sooner if the reaction wall and RNA continue to be effective. Model predictions suggest that the IM groundwater remedial goal of 1,000 µg/L for 1,1,1-TCA may not be attained for 95 years without source treatment.

If future LTM data indicate that 1,1,1-TCA discharge to Long-Branch Creek is a concern despite RNA and the reactive barrier walls, then Alternative 2 may be implemented as a contingency. Implementation of Alternative 2 will require further contaminant source delineation and characterization. Source delineation could be performed using minimally intrusive soil gas survey (passive or active) techniques to identify 1,1,1-TCA and CAH "hot spots" that may correspond to a NAPL source area. An active soil gas survey involves driving a perforated probe into the shallow subsurface and extracting soil gas using a vacuum pump. The gas sample can be analyzed onsite for the contaminant(s) of concern (COCs) using a gas chromatograph (GC). A potential advantage of an active survey is that variations in soil gas quality with depth can easily be investigated by collecting samples at multiple depths. Increases in contaminant concentrations with depth indicate the presence of a subsurface source as opposed to a surficial source.

With a passive soil gas survey, an absorbent collector is placed a few inches to a few feet below the ground surface for a period of time, and is then removed and analyzed onsite or at a fixed-base laboratory for the COCs. A potential advantage of a passive survey is that passive collectors tend to be more sensitive to subsurface contaminant concentrations because they are left in place over a period of time. Prior to performing a full-scale survey, both active and passive samples could be collected at a small number of locations and the results compared to determine the optimal method for the OU-4 area.

The estimated cost for a 150-point passive soil gas survey is approximately \$30,000. This cost includes a letter work plan, sampler installation/removal, laboratory analysis,

and a results letter report. If one or more CAH "hot spots" are identified from the passive soil gas survey, then drilling and soil/groundwater sampling would be required to further define subsurface physical and chemical conditions and costs would vary accordingly.

Under any scenario (Alternative 1 or 2 with reactive barrier walls), model results suggest that the drinking water criterion and aquatic life standard for 1,1,1-TCA will not be exceeded in Long Branch Creek at any time in the future. This conclusion is supported by the lack of quantifiable 1,1,1-TCA detections in Long Branch Creek. In addition, the shallow groundwater at the site and surface water in the on-Base reach of Long Branch Creek are not being used as drinking water or irrigation sources, and may not represent viable water sources due to the currently restricted nature of the Base. During and after source removal activities, if LTM results demonstrate that the plume is stable or receding and surface water standards are not exceeded, then the frequency of LTM could potentially be reduced (e.g., to once every 2 years or more).

### **SECTION 7**

### LONG-TERM MONITORING PLAN

### 7.1 OVERVIEW

In keeping with the requirements of the preferred remedial alternative for OU-4 (Section 7), a long-term groundwater monitoring plan was developed. The purpose of this component of the preferred remedial alternative for the site is to assess site conditions over time and the effectiveness of naturally occurring processes at reducing contaminant mass and minimizing contaminant migration, and evaluate the need for additional remediation.

The long-term groundwater monitoring plan for OU-4 presented in this section describes a monitoring scheme for the next 20 years for planning and budgeting purposes. It should be noted that this plan was devised on the basis of currently available data; the LTM program may be progressively revised as new data are obtained during this 20-year period. The LTM plan consists of identifying the locations of LTM wells and developing a groundwater sampling and analysis strategy to accomplish the following objectives:

- Monitor changes in site conditions, including plume magnitude and extent, over time;
- Evaluate the 1,1,1-TCA degradation rate and assess the effectiveness of naturally occurring processes at reducing contaminant mass and minimizing contaminant migration;
- Assess the need for additional remediation; and
- Verify the predictions of the numerical contaminant fate and transport model.

The strategy described in this section is designed to monitor plume migration over time and to verify that natural and engineered remedial processes are adequately protecting potential receptors. In the event that data collected under this LTM program indicate that naturally occurring processes are insufficient to adequately protect human health and the environment, contingency controls to more aggressively remediate the dissolved 1,1,1-TCA plume would be necessary.

### 7.2 MONITORING NETWORKS

Two separate types of wells will be used at the site as part of the LTM program. The first type consists of LTM wells to verify the results of the MODFLOW/MT3D modeling and to ensure that natural attenuation is occurring at rates sufficient to

adequately limit plume expansion and deleterious impact to surface water standards. The second type of well consists of a sentry well to verify that no 1,1,1-TCA exceeding IM groundwater quality guidelines migrate beyond the area under institutional control.

### 7.2.1 Groundwater Monitoring Network and Sampling Frequency

Annual LTM of three new LTM wells and 8 existing monitoring wells located within the CAH plume is recommended. The locations of the proposed and existing wells to be sampled are shown on Figure 7.1. Three of the proposed LTM wells are located along the approximate centerline of the east/southeast contaminant plume lobe to help monitor the progress of natural attenuation. These proposed LTM wells also were positioned to further delineate maximum plume concentrations, especially in the source area near MPC where no permanent well currently are located. The existing wells to be sampled include MW-105, MW-112, MW-113, MW-114, MW-115, MW-116, MW-117, and MW-118 (Figure 7.1). Well MW-116 is located upgradient from the majority of the plume, and the remaining wells are located within or transverse to the contaminant plume lobes. The transverse well locations are intended to monitor any potential expansion of the plume in the future. For scoping and costing purposes, this LTM plan assumes that all selected LTM wells initially will be sampled annually. Performance of annual sampling for 5 years (1998 through 2002) is recommended to build a historical water quality database for the site.

Following completion of the 5-year annual LTM period, biennial (every other year) LTM of all wells for an additional 15 years (2003 through 2018) is assumed. Estimates of the groundwater/CAH migration rate and direction should be progressively refined as new data are obtained to aid in plume definition and tracking. In addition, the LTM results can be compared to model-predicted concentrations to assess whether the numerical model has accurately simulated plume persistence over time.

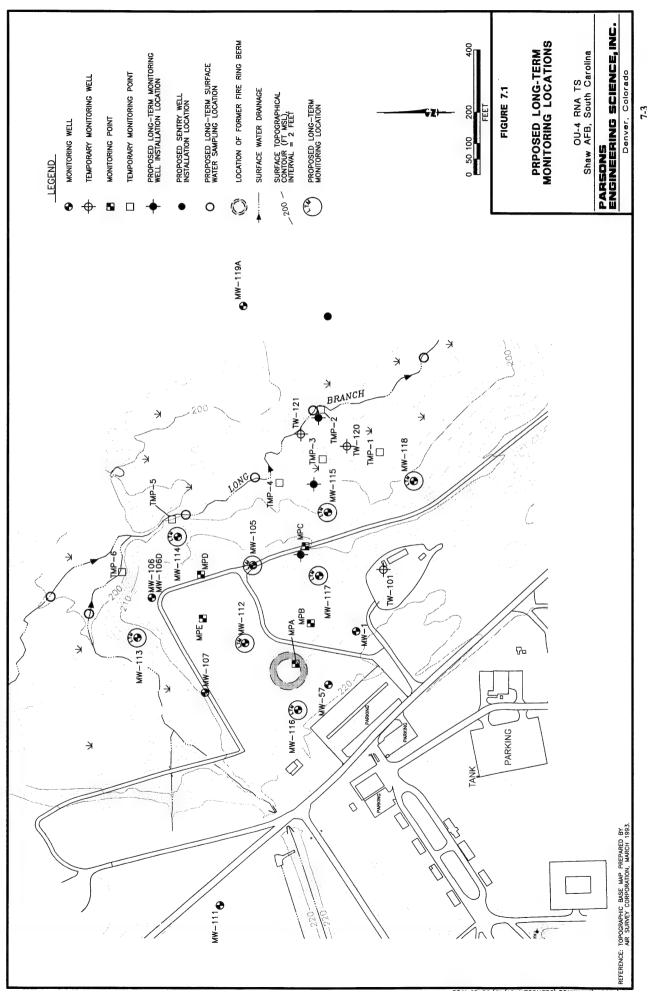
### 7.2.2 Sentry wells

The second type of monitoring well to be used at OU-4 is a sentry well. A single sentry well is proposed approximately 300 feet east of Long Branch Creek at the point of maximum contaminant discharge. The purpose of the sentry well is to confirm that the dissolved plume is not migrating beyond the area under institutional control. The preliminary sentry well location is shown on Figure 7.1.

### 7.3 SURFACE WATER MONITORING NETWORK AND SAMPLING FREQUENCY

Annual sampling of a minimum of six surface water stations during the 5-year annual LTM period described above is recommended. The six stations are located along Long Branch Creek upstream, within, and downstream from the reach projected to receive contaminant discharge (Figure 7.1). Data from these samples will be used to evaluate the impact of groundwater discharge on the quality of the surface water, and the effects of natural attenuation (e.g., dilution, volatilization) on contaminant concentrations (if any) in the creek.

The need for additional surface water quality monitoring following completion of the 5-year annual monitoring period should be assessed on the basis of the monitoring



results. Biennial monitoring of surface water for 15 years starting in year 2004 is assumed for costing purposes.

### 7.4 ANALYTICAL PROTOCOL

All LTM wells should be sampled and analyzed to determine compliance with chemical-specific remediation goals and to verify the effectiveness of remediation at the site. Groundwater level measurements should be made during each sampling event. Groundwater samples from LTM wells should be analyzed for the parameters listed in Table 7.1. Surface water samples should be analyzed for VOCs using USEPA Method 8260b or 8021b.

### 7.5 PERIODIC LTM PLAN REVIEW

The LTM plan should be periodically reviewed and revised as appropriate on the basis of available groundwater quality data. For example, if the sampling results obtained during the 15-year biennial sampling period demonstrate that the plume is in equilibrium (not receding or migrating downgradient) or is receding back toward the source area, then the sampling frequency could be reduced appropriately. If the data collected at any time during the monitoring period indicate the need for additional remedial activities at the site, sampling frequency should be adjusted accordingly. If sampling results indicate that geochemical conditions in the plume area are stable over time (e.g., nitrate, sulfate, and ferrous iron concentrations), then the sampling frequency for these parameters could be reduced.

# TABLE 7.1 LONG-TERM GROUNDWATER MONITORING ANALYTICAL PROTOCOL OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

	n Fixed-Base Laboratory	ss Field D,	w- Field	w- Field	w- Field	a Field sure ell	a Fixed-base
Sample Volume, Sample	Container, Sample Preservation	Collect 100 mL of water in a glass container; for Method A3500-FeD, acidify with hydrochloric acid per method	Measure at well-head using a flow- through cell	Measure at well-head using a flow-through cell	Measure at well-head using a flow- through cell	Collect 100-250 mL of water in a glass or plastic container or measure at wellhead using flow-through cell	Collect up to 40 mL of water in a glass or plastic container; cool to 4°C; analyze within 48 hours
Recommended	Frequency of Analysis	Each sampling event	Each sampling event	Each sampling event	Each sampling event	Each sampling event	Each sampling event
	Data Use	May indicate an anaerobic degradation process due to the depletion of oxygen, nitrate, and manganese	Purging adequacy; metabolism rates for microorganisms depend on temperature	Purging adequacy; concentrations less than 1 mg/L generally indicate an anaerobic pathway	Purging adequacy; aerobic and anaerobic processes are pH-sensitive	General water quality parameter used as a marker to verify that site samples are obtained from the same groundwater system	Substrate for microbial respiration if oxygen is depleted
	Comments	Filter if turbid	Measure at well- head	Measure at well-head; refer to Method A4500 for a comparable laboratory procedure	Measure at well-head	Measure at well- head	Method E300 is a Handbook <sup>20</sup> method; method SW9056 is an equivalent
	Method/Reference	Colorimetric A3500-Fe D or Hach 25140-25	E170.1, direct-reading meter	Dissolved oxygen meter	E150.1/SW9040, direct-reading meter	E120.1/SW9050, direct-reading meter	IC method E300 or method SW9056; colorimetric, method E353.2
	Analyte	Ferrous Iron (Fe <sup>2+</sup> )	Temperature**	Dissolved Oxygen**	**Hd	Conductivity**	Nitrate (NO <sub>3</sub> -1)

7-5

## TABLE 7.1 (Concluded) LONG-TERM GROUNDWATER MONITORING ANALYTICAL PROTOCOL SHAW AFB, SOUTH CAROLINA OU4 RNA TS

Analyte	Method/Reference	Comments	Data Use	Recommended Frequency of Analysis	Sample Volume, Sample Container, Sample Preservation	Field or Fixed-Base Laboratory
Sulfate (SO <sub>4</sub> <sup>2</sup> )	IC method E300 or method SW9056 or Hach SulfaVer 4 method	Method E300 is a Handbook method; method SW9056 is an equivalent procedure. Hach method is Photometric	Substrate for anaerobic microbial respiration	Each sampling event	Collect up to 40 mL of water in a glass or plastic container; cool to 4°C	Fixed-base or field (for Hach method)
Oxidation- Reduction Potential (ORP)**	A2580 B, direct-reading meter	Measurements are made with electrodes; results are displayed on a meter; samples should be protected from exposure to atmospheric oxygen	The redox potential of groundwater influences and is influenced by biologically mediated reactions; the redox potential of groundwater may range from more than 200 mV to less than -400 mV	Each sampling event	Measure at well-head using a flow- through cell	Field
Methane	RSKSOP-114 modified to analyze water samples for methane by headspace sampling with dual thermal conductivity and flame ionization detection.	Method published and used by the USEPA National Risk Management Research Laboratory	The presence of methane indicates the presence of sufficiently reducing conditions for reductive dehalogenation to occur	Each sampling event	Collect water samples in 40 mL volatile organic analysis (VOA) vials with butyl gray/Teflon-lined caps (zero headspace); cool to 4°C	Fixed-base
Aromatic Volatile Organics**	GC/MS method SW8260b or GC method SW8021b.	SW8260 is the Handbook method.	Measured for regulatory compliance.	Each sampling event	Collect water samples in a 40 mL VOA vial; cool to 4°C; add hydrochloric acid to pH < 2	Fixed-base

7-6

a/ Handbook refers to the AFCEE (1993) "Handbook for the Installation Restoration Program (IRP) Remedial Investigations and Feasibility Studies (RI/FS)."

Note: \*\* indicates parameters that should be monitored in both LTM and sentry wells; all other parameters should only be monitored in LTM wells.

### **SECTION 8**

### CONCLUSIONS AND RECOMMENDATIONS

This report presents the results of a TS conducted to evaluate the use of natural attenuation for remediation of CAH-contaminated groundwater at OU-4 at Shaw AFB, South Carolina. The finite-difference models MODFLOW and MT3D<sup>96®</sup> were used in conjunction with site-specific geologic, hydrologic, and laboratory analytical data to simulate the migration and biodegradation of CAHs dissolved in groundwater. To obtain the data necessary for the RNA demonstration, soil, groundwater, surface water, and sediment samples were collected from the site and analyzed. Physical and chemical data collected under this program were supplemented (where necessary) with data collected during previous site characterization events.

Comparison of CAH, TOC, electron acceptor, and byproduct data provides strong evidence that CAHs dissolved in groundwater at OU-4 are being naturally degraded. CAH compounds such as PCE, TCE, and 1,1,1-TCA are undergoing biologically facilitated (anaerobic) reductive dechlorination (e.g., TCE to DCE or 1,1,1-TCA to 1,1-DCA) and abiotic losses through dehydrohalogenation (e.g., 1,1,1-TCA to 1,1-The effects of these anaerobic processes, combined with physical dilution and chemical sorption, is causing the parent solvent concentrations at the site to decrease by approximately 2 orders-of-magnitude between the source area and Long Branch Creek. The anaerobic destruction of DCE and DCA (the byproducts of TCE and TCA dechlorination) also is likely occurring in the more anaerobic portion of the groundwater plume near the source area; however, DCE and DCA are reduced to a lesser extent between the source area and Long Branch Creek than observed with TCE and TCA. Near the leading edge of the plume near the creek, reductive dechlorination appears to be less prevalent, and compounds such as DCE, DCA, VC, and the chlorobenzenes likely are aerobically degraded or diluted. These compounds either were not detected or were detected at very low concentration in Long Branch Creek.

Site-specific hydrogeologic and laboratory analytical data were used in the numerical groundwater model to simulate the effects of advection, dispersion, sorption, and biodegradation on the fate and transport of the total dissolved CAH plume and the 1,1,1-TCA plume. A surrogate IM remedial groundwater standard for 1,1,1-TCA was defined by SCDHEC (1997), and this compound is used as an indicator compound in this report. Therefore, the fate and transport of 1,1,1-TCA was simulated using the numerical model. During model calibration, model parameters that could not be obtained from existing site data were estimated using common literature values for aquifer materials similar to those found at the site. Model predictions were found to be very sensitive to the magnitude of the first-order CAH decay rate assigned to the shallow portion of the surface aquifer at OU-4.

Remedial Alternative 1 incorporates natural attenuation, but does not include engineered remediation. The model used to simulate this alternative (OU4MODA) incorporates the assumption that dissolution from mobile and/or residual NAPL in the source area into groundwater will naturally decrease at a geometric rate of 3 percent per year (each concentration was decreased by a factor equal to 3 percent of the previous year's concentration) during the 100-year predictive period. The results of the model indicate that CAH concentrations will continue migrating from the source area to the edge of Long Branch Creek and are not predicted to decrease below the IM goal for 1,1,1-TCA (the surrogate compound for the site) of 1,000  $\mu$ g/L for 95 years. CAH concentrations discharging to Long Branch Creek are significantly lower than source area concentrations, and appear to have a minimal impact on surface water quality based on the lack of 1,1,1-TCA detections in the creek.

Model OU4MODB, used to simulate remedial Alternative 2, assumes that partial source removal via application of an engineered remedial technology such as SVE would reduce total CAH and 1,1,1-TCA dissolution from NAPL within a 5-year operational period. This model incorporates the assumption that the SVE system will reduce the dissolution of CAH compounds into groundwater by 8 percent per year for the first 2 years, 5 percent per year for the next 3 years, and 3 percent per year for the remainder of the 100 year model simulation. Results of these models indicate that source reduction at the assumed rates would result in only marginal decreases in dissolved CAH and 1,1,1-TCA concentrations within the plume relative to Alternative 1. Model OU4MODB predicts that leachable CAH contamination will be reduced by approximately 12 percent by calendar year 2008 (5 years of SVE followed by 5 years of natural weathering) over OU4MODA simulations and that 1,1,1-TCA concentrations will decrease below 1,000 μg/L within approximately 90 years.

A third groundwater model was used to simulate the potential effects of a groundwater extraction and treatment system at the site. Model OU4MODC was identical to OU4MODA, except that it used a single pumping well placed in the currently delineated 1,1,1-TCA plume source, near monitoring point MPC. The results of this model indicate that groundwater extraction and treatment at rates above 20 gpm will likely mitigate any contact of the 1,1,1-TCA plume lobe with Long Branch Creek. However, long-term groundwater extraction and treatment is not the recommended remedial method for the site. The permeable reactive barrier wall installed in November 1998 is preferable to groundwater extraction and treatment and may degrade groundwater contaminants before entry into Long Branch Creek. Future groundwater sampling will determine the effectiveness of the barrier wall and if LTM period or sampling frequency can be modified. Any groundwater treatment or interception system would potentially continue indefinitely unless the source area were remediated simultaneously. Furthermore, groundwater extraction and treatment would alter the hydraulic and geochemical characteristics of the site, thereby affecting contaminant decay rates and preventing the collection of meaningful data for monitored natural attenuation.

On the basis of this evaluation, Alternative 2, which combines natural attenuation with NAPL source removal through SVE, is recommended. Implementation of Alternative 2 will reduce the overall remediation time. A properly designed SVE system may achieve removal rates greater than assumed in the model simulations, further accelerating remediation rates at OU-4. Prior to installation of the SVE system,

the performance of a minimally intrusive soil gas survey should be considered to pinpoint subsurface CAH and 1,1,1-TCA "hot spots." A simple active or passive soil-gas survey would aid in identifying the location of mobile or residual LNAPL, thereby refining the design and effectiveness of any engineered remediation system.

The beneficial effects of natural attenuation are included in Alternative 2. Available data indicate that the total CAH and 1,1,1-TCA plumes are stabilized and biodegrading east of the source area. The degree to which RNA will continue to prevent significant downgradient migration of the plume is not known with certainty, but will become more apparent as successive LTM sampling events are performed. In any event, model results suggest that the drinking water criterion and aquatic life standard for 1,1,1-TCA will not be exceeded in Long Branch Creek at any time in the future; this observation is supported by the lack of quantifiable 1,1,1-TCA detections in Long Branch Creek. Shallow groundwater at the site or surface water in Long Branch Creek are not being used as a drinking water or irrigation source, and may not represent a viable potable water source due to the proximity and restricted nature of the base.

To assess the effectiveness of both naturally-occurring processes and SVE at reducing contaminant mass and minimizing contaminant migration, groundwater from a minimum of 8 existing monitoring wells and 4 proposed monitoring wells should be sampled annually for 5 years to build a historical groundwater quality database for the site. Six surface water stations also should be sampled annually during this 5-year period. Following completion of the 5-year annual LTM period, the LTM program should be reassessed. If the annual sampling results indicate that temporal changes in plume magnitude and extent could be adequately monitored by less-frequent sampling events, then the frequency of LTM could be reduced appropriately. The number and location of wells and surface water stations selected for continued monitoring should be determined on the basis of results from the initial, 5-year LTM period.

The LTM plan should be periodically reviewed and revised as appropriate on the basis of available groundwater quality data. For example, if sampling results demonstrate that the plume is in equilibrium (not receding or migrating downgradient) or is receding toward the source area, then the sampling frequency could be reduced appropriately. If the data collected at any time during the monitoring period indicate the need for additional remedial activities at the site, sampling frequency should be adjusted accordingly and the appropriate remedial actions (e.g., air sparging, iron filing trench as briefly described in Section 6.2.4) should be evaluated and implemented.

### **SECTION 9**

### REFERENCES

- Abriola, L.M., 1996, Organic liquid contaminant entrapment and persistence in the subsurface: Interphase mass transfer limitation and implications for remediation. 1996 Darcy Lecture, National Ground Water Association, presented at Colorado School of Mines, October 25, 1996.
- Altenschmidt, U. and Fuchs, G., 1991, Anaerobic degradation of toluene in denitrifying Pseudomonas sp.: Indication for toluene methylhydroxylation and benzoyl-CoA as central aromatic intermediate: Arch. Microbial., vol. 156, p. 152-158.
- Alvarez-Cohen, L.M. and McCarty, P.L., 1991a, Effects of toxicity, aeration, and reductant supply on trichloroethylene transformation by a mixed methanotrophic culture. *Appl. Environ. Microbiol.*, 57(1):228-235.
- Alvarez-Cohen, L.M. and McCarty, P.L., 1991b, Product toxicity and cometabolic competitive inhibition modeling of chloroform and trichloroethylene transformation by methanotrophic resting cells. *Appl. Environ. Microbiol.*, 57(4):1031-1037.
- Alvarez, P.J.J., and Vogel, T.M., 1991, Substrate interactions of benzene, toluene, and para-xylene during microbial degradation by pure cultures and mixed culture aquifer slurries: Applied Environmental Microbiology, vol. 57, p. 2981-2985.
- Arciero, D., Vannelli, T., Logan, M., and Hooper, A.B., 1989, Degradation of trichloroethylene by the ammonia-oxidizing bacterium *Nitrosomonas europaea*. *Biochem. Biophys. Res. Commun.*, 159:640-643.
- Ashworth, R.A., Howe, G.B., Mullins, M.E., and Rogers, T.N., 1988, Air-water partitioning coefficients of organics in dilute aqueous solutions. *Journal of Hazardous Materials*, 18:25-36.
- Atlas, R.M., 1981, Microbial degradation of petroleum hydrocarbons an environmental perspective: Microbiological Reviews, vol. 45, no. 1, p. 180-209.
- Atlas, R.M., 1984, Petroleum Microbiology: Macmillan Publishing Company, New York.
- Atlas, R.M., 1988, Microbiology Fundamentals and Applications: Macmillan Publishing Company, New York.
- Baedecker, M.J., Siegel, D.I., Bennett, P.C., Cozzarelli, I.M., 1988, The fate and effects of crude oil in a shallow aquifer: I. The distribution of chemical species

- and geochemical facies, In: G.E. Mallard, and S.E. Ragone, editors, U.S. Geological Survey Toxic Substances Hydrology Program, Proceedings of the Technical Meeting, Phoenix, Arizona, September 26-30, 1988: U.S. Geological Survey Water-Resources Investigations Report 88-42320, p. 29-22.
- Baedecker, M.J., and Cozzarelli, I.M., 1991, Geochemical modeling of organic degradation reactions in an Aquifer contaminated with Crude Oil: U.S. Geological Survey Water-Resources Investigations Report 91-4034. Reston, VA. p 627-632.
- Ball, H.A., Reinhard, M., and McCarty, P.L., 1991, Biotransformation of monoaromatic hydrocarbons under anoxic conditions, In: In Situ Bioreclamation, Applications and Investigations for Hydrocarbon and Contaminated Site Remediation. Eds. R.E. Hinchee and R.F. Olfenbuttel. Butterworth-Heinemann. Boston, MA. p. 458-463.
- Barker, J.F., Patrick, G.C., and Major, D., 1987, Natural attenuation of aromatic hydrocarbons in a shallow sand aquifer: Ground Water Monitoring Review, Winter, 1987, p. 64-71.
- Bartha, R., 1986, Biotechnology of petroleum pollutant biodegradation: Microbial Ecology, vol. 12, p. 155-172.
- Bauman, B., 1991, Biodegradation research of the American Petroleum Institute. Presented at: In Situ Bioreclamation: Application and Investigation for Hydrocarbons and Contaminated Site Remediation. San Diego, CA. March 19-21, 1991.
- Bear, J., 1979, *Hydraulics of Groundwater*. McGraw-Hill International Book Co., New York.
- Beller, H.R. Reinhard, M., and Grbic-Galic, D., 1992, Metabolic byproducts of anaerobic toluene degradation by sulfate-reducing enrichment cultures: Appl. Environ. Microbiol., vol. 58, p. 3192-3195.
- Borden, R.C., 1991, Simulation of enhanced in situ biorestoration of petroleum hydrocarbons. In: In Situ bioreclamation: Application and Investigation for Hydrocarbons and contaminated Site Remediation. Eds., R.E. Hinchee and R. F. Olfenbuttel. Butterworth-Heinemann. p. 529-534.
- Bouwer, E.J., 1992, Bioremediation of Subsurface Contaminants. In: Mitchell, R.(ed.), *Environmental Microbiology*. Wiley-Liss, New York.
- Bouwer, E.J., 1994, Bioremediation of chlorinated solvents using alternate electron acceptors. In: Norris, R.D., Hinchee, R.E., Brown, R., McCarty, P.L., Semprini, L., Wilson, J.T., Kampbell, D.H., Reinhard, M., Bouwer, E.J., Borden, R.C., Vogel, T.M., Thomas, J.M., and Ward, C.H. (eds.), Handbook of Bioremediation. Lewis Publishers.
- Bouwer, E.J., Rittman, B.E., and McCarty, P.L., 1981, Anaerobic degradation of halogenated 1- and 2-carbon organic compounds. *Environ. Sci. Technol.*, 15(5):596-599.

- Bouwer, E.J. and Wright, J.P., 1988, Transformations of trace halogenated aliphatics in anoxic biofilm columns. *J. Contam. Hydrol.*, 2:155-169.
- Bouwer, H., 1989, The Bouwer and Rice slug test an update. *Ground Water*, 27(3):304-309.
- Bouwer, H. and Rice, R.C., 1976, A slug test for determining hydraulic conductivity of unconfined aquifers with completely or partially penetrating wells. *Water Resources Research*, 12(3):423-428.
- Bradley, P.M. and Chapelle, F.H., 1996, Anaerobic mineralization of vinyl chloride in Fe(III)-reducing aquifer sediments. Accepted for publication in: *Environmental Science and Technology*.
- Brown, R.A., Dey, J.C. and McFarland, W.E., 1991, Integrated site remediation combining groundwater treatment, soil vapor extraction, and bioremediation, In: In Situ Bioreclamation: Application and Investigation for Hydrocarbons and Contaminated Site Remediation. Eds., R.E. Hinchee and R. F. Olfenbuttel. Butterworth-Heinemann. p. 444-449.
- Buscheck, T.E. and Alcantar, C.M., 1995, Regression techniques and analytical solutions to demonstrate intrinsic bioremediation. In: *Proceedings of the 1995 Battelle International Conference on In-Situ and On Site Bioreclamation*. April.
- Butler, B.J., and Barker, J.F., 1996, Chemical and microbiological transformation and degradation of chlorinated solvent compounds, <u>In</u>, Pankow, J.F., and Cherry, J.A., eds., Dense Chlorinated Solvents and Other DNAPLs in Groundwater: History, Behavior, and Remediation: Waterloo Press, Waterloo, Ontario, p. 267-312.
- Chapelle, F.H., 1993, Ground-Water Microbiology and Geochemistry. John Wiley & Sons, Inc., New York.
- Chapelle, F.H., McMahon, P.B., Dubrovsky, N.M., Fujii, R.F., Oaksford, E.T., and Vroblesky, D.A., 1995, Deducing the distribution of terminal electron-accepting processes in hydrologically diverse groundwater systems. Water Resources Research, 31:359-371.
- Chiang, C.Y., Salanitro, H.P., Chai, E.Y., Colthart, H.D., and Klein, C.L., 1989, Aerobic biodegradation of benzene, toluene, and xylene in a sandy aquifer data analysis and computer modeling: Ground Water, vol. 27, no. 6, p. 823-834.
- Cline, P.V. and Delfino, J.J., 1989, Transformation kinetics of 1,1,1-trichloroethane to the stable product 1,1-dichloroethene. In: *Biohazards of Drinking Water Treatment*. Lewis Publishers, Inc, Chelsea, MI.
- Cohen, R. M., and J. W. Mercer, 1993, *DNAPL Site Evaluation*. CRC Press, Inc., Boca Raton, Florida.
- Cozzarelli, I.M., Eganhouse, R.P., and Baedecker, M.J., 1990, Transformation of monoaromatic hydrocarbons to organic acids in anoxic ground water environment: Environmental and Geological Water Science, vol. 16.

- Criddle, C.S., J.T. DeWitt, and P.L. McCarty, 1990, Reductive dehalogenation of carbon tetrachloride by Escherichia coli K-12. Applied and Environmental Microbiology, 56:3247-3254.
- Davis, J.W. and Carpenter, C.L., 1990, Aerobic biodegradation of vinyl chloride in groundwater samples. *Applied and Environmental Microbiology*, 56:3878.
- Davis, J.W., Klier, N.J., and Carpenter, 1994, Natural biological attenuation of benzene in ground water beneath a manufacturing facility: Ground Water, vol. 32, no. 2, p. 215-226.
- de Bont, J.A.M., Vorage, M.J.W., Hartmans, S., and van den Tweel, W.J.J., 1986, Microbial degradation of 1,3-dichlorobenzene: Applied and Environmental Microbiology, v. 52, p. 677-680.
- De Bruin, W.P., Kotterman, M.J.J., Posthumus, M.A., Schraa, G., and Zehnder, A.J.B., 1992, Complete biological reductive transformation of tetrachloroethene to ethane: Applied and Environmental Microbiology, v. 58, no. 6, p. 1966 2000.
- DeStefano, T.D., Gossett, J.M., and Zinder, S.H., 1991, Reductive dehalogenation of high concentrations of tetrachloroethene to ethene by an anaerobic enrichment culture in the absence of methanogenesis. *Appl. Environ. Microbiol.*, 57(8):2287-2292.
- Dewulf, J., Drijvers, D., and Van Langenhove, H., 1995, Measurement of Henry's law constant as function of temperature and salinity for the low temperature range. *Atmos. Environ.*, 29(3):323-331.
- Domenico, P.A., and Schwartz, F.W., 1990, Physical and Chemical Hydrogeology. John Wiley and Sons, New York, New York, 824p.
- Edwards, E., Wills, L.E., Grbic-Galic, D., and Reinhard, M., 1991, Anaerobic degradation of toluene and xylene--evidence for sulfate as the terminal electron acceptor, In: In Situ Bioreclamation, Applications and Investigations for Hydrocarbon and Contaminated Site Remediation. Eds. R.E. Hinchee and R.F. Olfenbuttel. Butterworth-Heinemann. Boston, MA. p. 463-471.
- Edwards, E.A., and Grbic-Galic, D., 1992, Complete mineralization of benzene by aquifer microorganisms under strictly anaerobic conditions: Applied and Environmental Microbiology, vol. 58, p. 2663-2666.
- Edwards, E.A., and Grbic-Galic, D., 1994, Anaerobic degradation of toluene and oxylene by a methanogenic consortium. Applied and Environmental Microbiology, v.60, p313-322.
- Engineering-Science, Inc., 1983, Installation Restoration Program, Phase I: Records Search, Shaw AFB, South Carolina.
- Feenstra, S. and Guiguer, N., 1996, Dissolution of dense non-aqueous phase liquids in the subsurface. In: Pankow, J.F., and Cherry, J.A. (eds.), *Dense Chlorinated Solvents And Other DNAPLS In Groundwater*. Waterloo Press, Portland, OR.

- Florida Department of Environmental Protection (FDEP), 1993, Primary and Secondary Drinking Water Standards, Orlando, Florida.
- Focht, R., Vogan, J., and S. O'Hannersin, 1996, Field application of reactive iron walls for *in-situ* degradation of volatile organic compounds in groundwater. *Remediation*, Summer, John Wiley & Sons, Inc.
- Folsom, B.R., Chapman, P.J., and Pritchard, P.H., 1990, Phenol and trichloroethylene degradation by *Pseudomonsa cepacia* G4: Kinetics and interactions between substrates. *Appl. Environ. Microbiol.*, 56(5):1279-1285.
- Franke, O.L., Reilly, T.E., and Bennett, G.D., 1987, Definition of boundary and initial conditions in the analysis of saturated ground-water flow systems an introduction. In: *United States Geological Survey Techniques of Water-Resources Investigations*, Book 3, Chapter B5.
- Freedman, D.L. and Gossett, J.M., 1989, Biological reductive dehalogenation of tetrachloroethylene and trichloroethylene to ethylene under methanogenic conditions. *Appl. Environ. Microbiol.*, 55(4):1009-1014.
- Geraghty & Miller Modeling Group, 1994, AQTESOLV Aquifer Test Solver, Version 2.0. Millersville, Maryland. October.
- Gibson, D.T., and Subramanian, V., 1984, Microbial degradation of aromatic hydrocarbons, In: Gibson, D.T., ed: Microbial Degradation of Organic Compounds, Marcel Dekker, Inc., p. 181-252.
- Godsey, E.M., 1994, Microbiological and geochemical degradation processes. Symposium on Intrinsic Bioremediation in Ground Water, Denver, CO. August 30 - September 1.
- Goldstein, R.M., Mallory, L.M., and Alexander, M., 1985, Reasons for Possible Failure of Inoculation to Enhance Biodegradation: Applied Environmental Microbiology, vol. 50, no. 4, p. 977-983.
- Grbic'-Galic', D., 1990, Anaerobic microbial transformation of nonoxygenated aromatic and alicyclic compounds in soil, subsurface, and freshwater sediments. In: Bollag, J.M. and Stotzky, G.(eds.), *Soil Biochemistry*. Marcel Dekker, Inc., New York.
- Haag, F., Reinhard, M., and McCarty, P.L., 1991, Degradation of toluene and p-xylene in an anaerobic microcosms: Evidence for sulfate as a terminal electron acceptor: Environ. Toxicol. Chem., vol. 10, p. 1379-1389.
- Haigler, B.E., Nishino, S.F., and Spain, J.C., 1988, Degradation of 1,2-dichlorobenzene by a *Pseudomonas sp.*: Applied and Environmental Microbiology, v. 54, p. 294-301.
- Harker, A.R. and Kim, Y., 1990, Trichloroethylene degradation by two independent aromatic-degrading pathways in *Alcaligenes eutrophus JMP134*. *Appl. Environ. Microbiol.*, 56(4):1179-1181.

- Hartmans, S. and de Bont, J.A.M., 1992, Aerobic vinyl chloride metabolism in *Mycobacterium aurum Li. Appl. Environ. Microbiol.*, 58(4):1220-1226.
- Henry, S.M., 1991, Transformation of trichloroethylene by methanotrophs from a groundwater aquifer. Ph.D. Thesis, Stanford University. Palo Alto, California.
- Hine, J., and Mookerjee, P.K., 1975, The Intrinsic Hydrophilic Character of Organic Compounds. Correlations in Terms of Structural Contributions: Journal or Organic Chemistry, 40(3):292-298.
- Hunter-Smith, R.J., Balls, P.W., and P.S. Liss, 1983, Henry's Law Constants and the Air-Sea Exchange of Various Low Molecular Weight Halocarbon Gases. *Tellus*, 35B:170-176.
- Hutchins, S.R., and Wilson, J.T., 1991, Laboratory and field studies on BTEX biodegradation in a fuel-contaminated aquifer under denitrifying conditions, In: In Situ Bioreclamation, Applications and Investigations for Hydrocarbon and Contaminated Site Remediation. Eds. R.E. Hinchee and R.F. Olfenbuttel. Butterworth-Heinemann, Boston, MA. p. 157-172.
- Hutchins, S.R., Sewell, G.W., Sewell, D.A., Kovacs, D.A., and Smith, G.A., 1991a, Biodegradation of monoaromatic hydrocarbons by aquifer microorganisms using oxygen, nitrate, or nitrous oxide as the terminal electron acceptor: Environ. Sci. Technol., vol. 25, no. 1, p. 68-76.
- Hutchins, S.R., Downs, W.C., Smith, G.B., Smith, J.T., Wilson, D.J., Hendrix, D.J., Fine, D.D., Kovacs, D.A., Douglass, R.H., and Blaha, F.A., 1991b, Nitrate for Biorestoration of an Aquifer Contaminated with Jet Fuel. U.S. Environmental Protection Agency. Robert S. Kerr Environmental Research Laboratory. Ada, Oklahoma. EPA/600/2-91/009. April, 1991.
- Jamison, V.W., Raymond, R.L., and Hudson, J.O. Jr., 1975, Biodegradation of highoctane gasoline in ground water: Developments in Industrial Microbiology, vol. 16.
- Jeffers, P.M., Ward, L.M., Woytowitch, L.M., and Wolfe, N.L., 1989, Homogeneous hydrolysis rate constants for selected chlorinated methanes, ethanes, ethenes, and propanes: Environmental Science and Technology, v. 23, p. 965-969.
- IT Corporation, 1997, Draft Chemical Oxidation Pilot Test Report, Operable Unit 4, IRP Site Number FT-01, Shaw AFB, SC, Contract No. DACW 45-93-D-0044, TERC Region 2, Delivery Order 0004, AF Project No. VLSB-95-7002, February, 1997.
- Klier, N.J., Welt, R.J., and Donberg, P.A., 1996, Aerobic biodegradation of dichlorethylenes in surface and subsurface soils. Submitted to *Chemosphere* by The Dow Chemical Company, Environmental Toxicology and Chemistry Laboratory, Health and Environmental Sciences. May.
- Konikow, L.F., and Bredehoeft, J.D., 1978, Computer model of two-dimensional solute transport and dispersion in groundwater. *United States Geological Survey, Techniques of Water Resources Investigations of the United States Geological Survey*, Book 7, Chapter C2, 90 p.

- Kuehne, D., and Buscheck, T., 1996, Survey of California marketing sites and analysis of monitoring well data: Chevron Research and Technology, unpublished report.
- Law Environmental, Inc., 1989, Remedial Action Plan, Shaw Air Force Base, July.
- Law Environmental, Inc., 1991, Draft Remedial Investigation and Feasibility Study Report for Site 1 Former Fire Training Area No. 1, Shaw AFB, Sumter, SC. Prepared for the US Army Corps of Engineers-Omaha Divisions, Decrease. December.
- Leahy, J.G., and Colewell, R.R., 1990, Microbial degradation of hydrocarbons in the environment: Microbiolgical Reviews, vol. 53, no. 3, p. 305-315.
- Lee, M.D., 1988, Biorestoration of Aquifers Contaminated with Organic Compounds: CRC Critical Reviews in Environmental Control. vol. 18. p. 29-89.
- Little, C.D., Palumbo, A.V., Herbes, S.E., Lidstrom, M.E., Tyndall, R.L., and Gilmer, P.J., 1988, Trichloroethylene biodegradation by a methane-oxidizing bacterium. *Appl. Environ. Microbiol.*, 54(4):951-956.
- Lovley, D.R., Chapelle, F.H., and Woodward, J.C., 1994, Use of dissolved H<sub>2</sub> concentrations to determine distribution of microbially catalyzed redox reactions in anoxic groundwater. *Environmental Science and Technology*, 28(7):1205-1210.
- Lovley, D.R. and Goodwin, S., 1988, Hydrogen concentrations as an indicator of the predominant terminal electron-accepting reaction in aquatic sediments. *Geochimica et Cosmochimica Acta*, 52:2993-3003.
- Lovely, D.R. and Phillips, E.J.P., 1988. Novel mode of microbial energy metabolism: Organic carbon oxidation coupled to dissimilatory reduction of iron or manganese. *Applied and Environmental Microbiology*, 54(6):1472-1480.
- Lovely, D.R. and Phillips, E.J.P., 1991, Enzymatic versus nonenzymatic mechanisms for Fe(III) reduction in aquatic sediments. *Environmental Science and Technology*, 26(6):1062-1067.
- Lyman, W.J., Reidy, P.J., and Levy, B., 1992, Mobility and Degradation of Organic Contaminants in Subsurface Environments. C.K. Smoley, Inc., Chelsea, MI.
- Mace, R.E., Fisher, R.S., Welch, D.M., and Parra, S.P., 1997, Extent, Mass, and Duration of Hydrocarbon Plumes from Leaking Petroleum Storage Tank Sites in Texas: Texas Bureau of Economic Geology, Geological Circular 97-1.
- Malone, D.R., Kao, C.M., and Borden, R.C., 1993, Dissolution and biorestoration of nonaqueous phase hydrocarbons models development and laboratory evaluation: Water Resources Research, vol. 29, no. 7, p. 2003-2213.
- Mayer, K.P., Grbic-Galic, D., Semprini, L., and McCarty, P.L., 1988, Degradation of trichloroethylene by methanotrophic bacteria in a laboratory column of saturated aquifer material. *Wat. Sci. Tech.* (Great Britain), 20(11/12):175-178.

- McCarty, P.L., Roberts, P.V., Reinhard, M., and Hopkins, G., 1992, Movement and transformations of halogenated aliphatic compounds in natural systems. In: Schnoor, J.L. (Ed.), Fate of Pesticides and Chemicals in the Environment. John Wiley & Sons, Inc., New York, New York.
- McCarty, P.L. and Semprini, L., 1994, Ground-water treatment for chlorinated solvents. In: *Handbook of Bioremediation*. Lewis Publishers, Boca Raton, FL.
- McDonald, G. and Harbaugh, A.W., 1988, A modular three-dimensional finite-difference groundwater flow model. US Geological Survey Techniques of Water Resources Investigations, Book 6, Chapter A1.
- Miller, R.E. and Guengerich, F.P., 1982, Oxidation of trichloroethylene by liver microsomal cytochrome P-450: Evidence for chlorine migration in a transition state not involving trichloroethylene oxide. *Biochemistry*, 21:1090-1097.
- Montgomery, J.J. 1996. Groundwater Chemicals Desk Reference. Lewis Publishers, Boca Raton.
- Moutoux, D.E., Benson, L.A., Swanson, T.H., Wiedemeier, T.H., Lenhart, J., Wilson, J.T., and Hansen J.E., 1996, Estimating the changing rate of anaerobic reductive dechlorination of chlorinated aliphatic hydrocarbons in the presence of petroleum hydrocarbons. *In: Proceedings of the 1996 API/NGWA Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water*. Houston, Texas. November.
- Murray, W.D. and Richardson, M., 1993, Progress toward the biological treatment of C<sub>1</sub> and C<sub>2</sub> halogenated hydrocarbons. Critical Reviews in Environmental Science and Technology, 23(3):195-217.
- Neely, W.B., and G.e. Blau, 1985, Environmental Exposure from Chemicals. Volume 1, CRC Press, Inc., Boca Raton, Florida.
- Nelson, M.J.K., Montgomery, S.O., O'Neille, E.J., and Pritchard, P.H., 1986, Aerobic metabolism of trichloroethylene by a bacterial isolate. *Appl. Environ. Microbiol.*, 52(2):949-954.
- Nishino, S.F., Spain, J.C., and Pettigrew, C.A., 1994, Biodegradation of chlorobenzene by indigeneous bacteria: Environ. Toxicol. Chem., v. 13, p. 871-877.
- Papadoulos, S.S., and Assoc., 1996, MT3D96: A Modular Three-Dimensional Transport Model for Simulation of Advection, Dispersion, and Chemical Reactions of Contaminants in Ground-Water Systems. Bethesda, Maryland.
- Parsons ES, 1994, Bioventing Test Work Plan For IRP Site FT-01 (Former Fire Training Area 1) and IRP Site SS-15 (POL Fuel Depot), Shaw AFB. May.
- Parsons ES, 1995, Draft Interim Bioventing Pilot Tests Results Report: IRP Site FT-01 (Former Fire Training Area 1) and IRP Site SS-15 (POL Fuel Depot), Shaw AFB, SC. January.

- Parsons ES, 1996, Memorandum to Mr. Rick Roller Re: Completion of Final Bioventing Test, Shaw AFB, Site FT-01 (Former Fire Training Area 1) and Site SS-15 (POL Fuel Depot). April.
- Parsons ES, 1997. Draft Work Plan for a Treatability Study in Support of Remediation by Natural Attenuation of Sites OU-4 (FT-1) and the Building 1613 Site.
- Reineke, W., and Knackmuss, H.-J., 1984, Microbial metabolism of haloaromatics: Isolation and properties of a chlorobenzene-degrading bacterium: European Journal of Applied Microbiology and Biotechnology, v. 47, p. 395-402.
- Reinhard, M., 1994, In-Situ bioremediation technologies for petroleum-derived hydrocarbons based on alternate electron acceptors (other than molecular oxygen). *Handbook of Bioremediation*. Lewis Publishers, Boca Raton, FL.
- Research Triangle Institute, 1986, Installation Restoration Program: Phase II: Confirmation 1 Quantification Stage 1, Shaw AFB, South Carolina, January.
- Rice, D.W., Grose, R.D., Michaelsen, J.C., Dooher, B.P., MacQueen, D.H., Cullen, S.J., Kastenberg, W.E., Everett, L.G., and Marino, M.A., 1995, California Leaking Underground Fuel Tank (LUFT) Historical Case Analyses: Environmental Protection Department, Environmental Restoration Division, Lawrence Berkely laboratories, UCRL-122207, prepared for the California State Water Resources Control Board.
- Rust E&I, 1993, Soil Vapor Extraction and Bioremediation Study Report, Operable Unit #4, Former Fire Training Area No. 1, Shaw AFB, SC. June.
- Rust E&I, 1994, Draft Final Site Inspection Report, Multi-Sites Group II, Shaw AFB, SC. January.
- Rust E&I, 1995, Final Remedial Investigation: Operable Unit #4, Former Fire Training Area No. 1 IRP Site No. FT-1, Shaw AFB, SC. February.
- Rust E&I, 1996, Final Feasibility Study/Corrective Measures Study Report, Proposal Plan, Operable Unit #4, Former Fire Training Area #1, IRP Site No. FT-1, SWMU #59.
- Sander, P., Wittaich, R.-M., Fortnagel, P., Wilkes, H., and Francke, W., 1991, Degradation of 1,2,4-trichloro- and 1,2,4,5-tetrachlorobenzene by *Pseudomonas* strains: Applied and Environmental Microbiology, v. 57, p. 1430-1440.
- Schraa, G., Boone, M.L., Jetten, M.S.M., van Neerven, A.R.W., Colberg, P.J., and Zehnder, A.J.B., 1986, Degradation of 1,2-dichlorobenzene by *Alcaligenes sp.* strain A175: Applied and Environmental Microbiology, v. 52, p. 1374-1381.
- SEC Donohue, Inc., 1992, Final Supplemental Remedial Investigation/Feasibility Study Work Plan: Operable Unit #4, Former Fire Training Area No. 1, Shaw Air Force Base, Sumpter, South Carolina.
- South Carolina Department of Health and Environmental Control (SCDHEC), July 14, 1997, Interim Measure (IM) letter to Mr. Randy Adams, 20 CES/CEV at Shaw AFB. Written by Mr. John T. Litton, Director, South Carolina DHEC, Division

- of Hazasrdous and Infectious Waste Management, Bureau of Land and Waste Management.
- Spain, J.C., 1996, Future vision: Compounds with potential for natural attenuation, In Proceedings of the Symposium on Natural Attenuation of Chlorinated Organics in Ground Water: EPA /540/R-96/509, Dallas TX, September 11-13, 1996.
- Spain, J., 1997, Personal communication between Jim Spain of the US Air Force Armstrong Laboratory and Matt Swanson of Parsons ES regarding potential anaerobic utilization of chlorobenzenes as substrates. March 7, 1997.
- Spain, J.C., and Nishino, S.F., 1987, Degradation of 1,4-dichlorobenzene by a *Pseudomonas sp.*: Applied and Environmental Microbiology, v. 53, p. 1010-1019.
- Spitz, K. and Moreno, J., 1996, A Practical Guide to Groundwater and Solute Transport Modeling. John Wiley & Sons, Inc., New York, 461 p.
- Stumm, W. and Morgan, J.J., 1981, Aquatic Chemistry. John Wiley & Sons, New York.
- Strack, O.D.L., 1989, Groundwater Mechanics: Prentice-Hall, Englewood Cliffs, New Jersey, 732 p.
- Suflita, J.M., and Townsend, G.T., 1995, The microbial ecology and physiology of aryl dehalogenation reactions and implications for bioremediation, In Young, L.Y., and Cerniglia, C.E., eds., Microbial Transformation and Degradation of Toxic Organic Chemicals: Wiley-Liss, New York, 654 p.
- Thierrin, J., Davis, G.B., Barber, C., Patterson, B.M., Pribac, F., Power, T.R., and Lambert, M., 1992, Natural degradation rates of BTEX compounds and naphthalene in a sulfate-reducing ground water environment, In: In-Situ Bioremediation Symposium "92". Niagara-on-the-Lake, Ontario, Canada. September 20-24, 1992: In Press.
- US Environmental Protection Agency (USEPA), 1988, Guidance for Conducting Remedial Investigations and Feasibility Studies (RI/FS) Under CERCLA. OSWER Directive 9355.3-01.
- USEPA, 1991, Water Quality Criteria Summary Concentrations, Office of Science and Technology, Health and Ecological Criteria Division, Ecological Risk Assessment Branch (WH-585), Human Risk Assessment Branch (WH-550D), Washington, D.C.
- USEPA, 1993, Revisions to the OMB Circular A-94 on Guidelines and Discount Rates for Benefit Cost Analysis, OSWER Directive 9355.3-20, June.
- van der Meer, J.R., Roelofsen, W., Schraa, G., and Zehnder, A.J.B., 1987, Degradation of low concentrations of dichlorobenzenes and 1,2,4-trichlorobenzene by *Pseudomonas* sp. strain P51 in nonsterile soil columns: FEMS Microbiology Letters, v. 45, p. 333-341.
- Vogel, T.M., 1994, Natural bioremediation of chlorinated solvents. In: *Handbook of Bioremediation*. Lewis Publishers, Boca Raton, FL.

- Vogel, T.M., Criddle, C.S., and McCarty, P. L., 1987, Transformations of halogenated aliphatic compounds. *Environmental Science and Technology*, 21(8):722-736.
- Vogel, T.M. and McCarty, P.L., 1985, Biotransformation of tetrachloroethylene to trichloroethylene, dichloroethylene, vinyl chloride, and carbon dioxide under methanogenic conditions. *Applied Environmental Microbiology*, 49(5):1080-1083.
- Vogel, T.M., Criddle, C.S., and McCarty, P.L., 1987, Transformations of halogenated aliphatic compounds: Environmental Science and Technology, vol. 21, no. 8, p. 722 736.
- Vroblesky, D.A., and Chapelle, F.H., 1994, Temporal and spatial changes of terminal electron-accepting processes in a petroleum hydrocarbon-contaminated aquifer and the significance for contaminant biodegradation: Water Resources Research, v. 30, no. 5, p. 1561-1570.
- Waterloo Hydrogeologic Software, 1996, Visual MODFLOW, Version 2.20. Waterloo, Ontario, Canada.
- Wiedemeier, T.H., Wilson, J.T., Kampbell, D.H., Miller, R.N., and Hansen, J.E., 1995, Technical Protocol For Implementing Intrinsic Remediation With Long-Term Monitoring For Natural Attenuation Of Fuel Contamination Dissolved In Groundwater. US Air Force Center for Environmental Excellence, San Antonio, Texas.
- Wiedemeier, T.H., Benson, L.A., Wilson, J.T., Kampbell, D.H., Hansen, J.E., and Miknis, R., 1996a, Patterns Of Natural Attenuation Of Chlorinated Aliphatic Hydrocarbons At Plattsburgh Air Force Base, New York. In: Conference on Intrinsic Remediation of Chlorinated Solvents. Salt Lake City, UT. April 2.
- Wiedemeier, T.H., Swanson, M.A., Moutoux, D.E., Gordon, E.K., Hansen, J.E., Haas, P., Wilson J.T., Kampbell, D.H., and Chapelle, F.H., 1996b, DRAFT *Protocol for Evaluating Natural Attenuation of Chlorinated Solvents in Groundwater*. Prepared for the Air Force Center for Environmental Excellence, Brooks Air Force Base, Texas. Revision 1.
- Wilson, J.T. and Wilson, B.H., 1985, Biotransformation of trichloroethylene in soil. *Appl. Environ. Microbiol.*, 49(1):242-243.
- Wilson, B.H., Smith, G.B., and Rees, J.F., 1986, Biotransformations of Selected Alkylbenzenes and Halogenated Aliphatic Hydrocarbons in Methanogenic Aquifer Material: A Microcosm Study: Environmental Science and Technology, 20(10):997-1002.
- Wilson, B.H., Bledsoe, B., and Kampbell, D., 1987, Biological processes occurring at an aviation gasoline spill site, In: R.C. Averett and D.M. McKnight editors, Chemical Quality of Water and the Hydrologic Cycle. Lewis Publishers, Chelsea, Michigan, p.125-137.

- Wilson, B.H., Wilson, J.T., Kampbell, D.H., Bledsoe, B.E., and Armstrong, J.M., 1990, Biotransformation of Monoaromatic and Chlorinated Hydrocarbons at an Aviation Gasoline Spill Site: Geomicrobiology Journal, 8:225-240.
- Wilson, J.T., 1996, Personal communication between John Wilson of the USEPA NRMRL and Todd Wiedemeier of Parsons ES regarding USEPA definition of the term "natural attenuation", citing USEPA guidance undergoing internal review.
- Young, L.Y., 1984, Anaerobic degradation of aromatic compounds, In: Gibson, D.R., ed.: Microbial Degradation of Aromatic Compounds. Marcel-Dekker, New York.
- Zheng, C., 1990, MT3D A Modular Three-Dimensional Transport Model for Simulation of Advection, Dispersion, and Chemical Reactions of Contaminants in Groundwater Systems; Prepared for the US Environmental Protection Agency Robert S. Kerr Environmental Research Laboratory, October 17, 1990.

### APPENDIX A

BOREHOLE LOGS, CPT/LIF RESULTS, MONITORING WELL CONSTRUCTION DIAGRAMS, MONITORING WELL/POINT SAMPLING FORMS, AQUIFER TEST RESULTS, AND SURVEY DATA.

2.5' Stick-up NETRATION-BLOWS PER FOOT 5 16 15 20 30 40 40 86 188 PEET DESCRIPTION ELEV. Firm, brown, silty sand (Fill) Firm, red and brown sandy silt (Fill) 5 Gray-brown silty sand with rock fragments and debris (Fill) 10 Very dense, light tan slightly silty sand SM 15 Very firm, gray sandy clay CL 20 Very dense, light tan, medium sand SP 25 30 Boring Terminated at 30' 35

TEST BORING RECORD

REMARKS:

2" PVC casing LOGGED BY M.Bond/ Elevation not surveyed CHECKED BY J.Wilson Water level measured 11/19/86

DRILLED BY R. Banks

LOGGED BY M. Bond/Babcock DATE STARTED

CHECKED BY J. Wilson DATE COMPLETE

BORING NUMBER PZ-101
DATE STARTED 11/8/86
DATE COMPLETED 11/8/86
602-2

21 Stick-up PENETRATION-BLOWS PER FOCT 5 10 15 26 30 40 50 80 100 PEET DESCRIPTION ELEV. Firm, brown and red-brown sand with asphalt and rock fragments Firm to very firm, gray to black medium sand with metallic debris 5 (Fill) 10 Dense, gray clayey sand SC 15 Very firm, gray fine to medium sand SW 20 Dense to very dense, white to light tan medium sand with some 25 gravel SW 30 Boring Terminated at 30'

TEST BORING RECORD

REMARKS:

2" PVC casing Elevation not surveyed Water level measured 11/19/86

DRILLED BY R.Banks
LOGGED BY M.Bond
CHECKED BY J.Wilson

BORING NUMBER PZ-102
DATE STARTED 11/11/86
DATE COMPLETED 11/11/86
602-2

TEST BORING RECORD 2' Stick-up PENETRATION-BLOWS PER FOOT 5 10 15 20 30 40 60 60 DEPTH DESCRIPTION ELEV. Loose to very firm red-brown to black, silty sand with asphalt and debris (Fill) 5 10 Dense, black asphaltic material (Fill) 15 Dense, gray and tan, medium sand SP 20 Dense to very dense, white sand with some gravel SW 25 30

REMARKS:

35

2" PVC casing

Water level measured 11/19/86 CHECKED BY J. Wilson

R. Banks DRILLED BY \_ M. Bond LOGGED BY

Boring Terminated at 35'

PZ-103 BORING NUMBER 11/11/87 DATE STARTED 11/11/87 DATE COMPLETED\_ 602-2\_\_ SERUN BOL

### TEST BORING RECORD

ELEV.	DEPTH FEET	DESCRIPTION	PENETRATION-BLOWS PER POOT 0 3 10 13 20 30 40 60 80 160
		Loose to firm; light tan sand SP	
	5	Gray clayey sand (Hydrocarbon smell) SC	
	10	Very firm, gray clayey sand SC	
	15		
	20	Very dense, white coarse sand SP	
	25	Boring Terminated at 25'	

REMARKS:

DRILLED BY R. Banks
LOGGED BY M. Bond
CHECKED BY J. Wilson

BORING NUMBER SB-111
DATE STARTED 11/7/86
DATE COMPLETED 602-2
JOB NUMBER

2.5' Stick-up PENETRATION-BLOWS PER POOT
3 10 13 20 30 40 40 80 100 PEET DESCRIPTION ELEV. Very firm, red-brown, silty sand with rock fragmenst (Fill) Firm, light tan and gray, slightly silty, fine to medium sand (Fill) 5 Firm, black, fine to medium sand with debris (Fill) 10 15 Very firm, gray, slightly silty, fine to medium sand SM Loose, light tan and gray, slightly silty fine sand SM 20 Firm to dense, white and tan 25 medium sand with some gravel SW 30 35 40

REMARKS:

Sheet 1 of 2 4" PVC casing Elevation not surveyed Water level measured 11/19/86

DRILLED BY R.Banks
LOGGED BY M.Bond
CHECKED BY J.Wilson

BORING NUMBER TW-101
DATE STARTED 11/9/86
DATE COMPLETED 602-2
JOB NUMBER

#### TEST BORING RECORD

ELEV.	DEPTH	DESCRIPTION		•	PEHE 3	TRATI	0 N- BL	30 4	-	007	۰ ،	••
	.45	Firm to dense, white and tan, medium sand with some gravel SW				·						
	50	Firm, white to gray, fine to medium sand SW										
		Very loose, white slightly clayey, fine to medium sand SC										
	-55										-	
	60	Boring Terminated at 60'										
		•										
			•									

REMARKS:

Sheet 2 cof 2

DRILLED BY R. Banks
LOGGED BY M. Bond
CHECKED BY J. Wilson

BORING NUMBER TW-101
DATE STARTED 11/9/86
DATE COMPLETED 11/9/86
602-2

From:

Kathleen Older

To:

internet.mcx: "jhansen@afceeb1.brooks.af.mil"

Date:

6/5/97 10:50am

Subject:

Wells at Shaw

SCAPS put in 5 power punch well points at Shaw. These well points have a mechanical packer above the screen. The screen intervals are approximately 10 feet and consist of 3 one meter long flush joint sections.

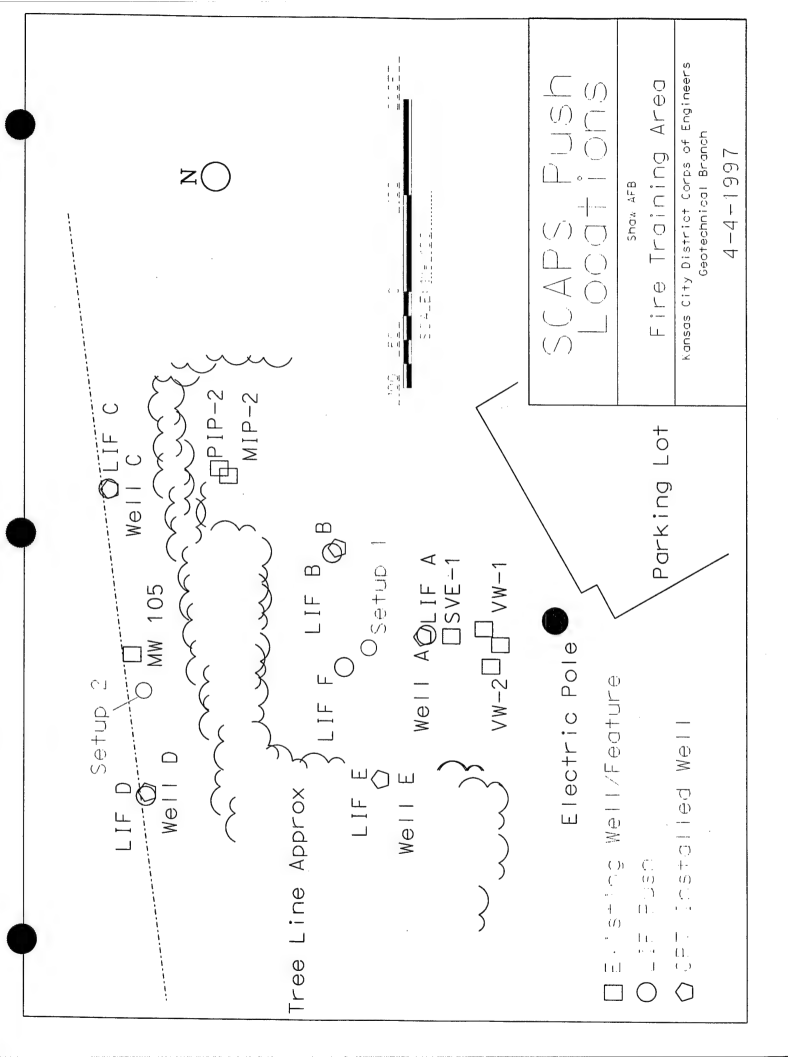
Well 5EFire is opened from 13 to 23 feet. water@ 13.0 ft Well at location B is opened 8 to 18 feet. water@ 13.5 ft Well at location A is opened 8 to 18 feet. water@ 12.9 ft Well at location D is opened 13 to 23 feet. water@ 15.1 ft Well at location C is opened 4 to 14 feet. water@ 9.9 ft

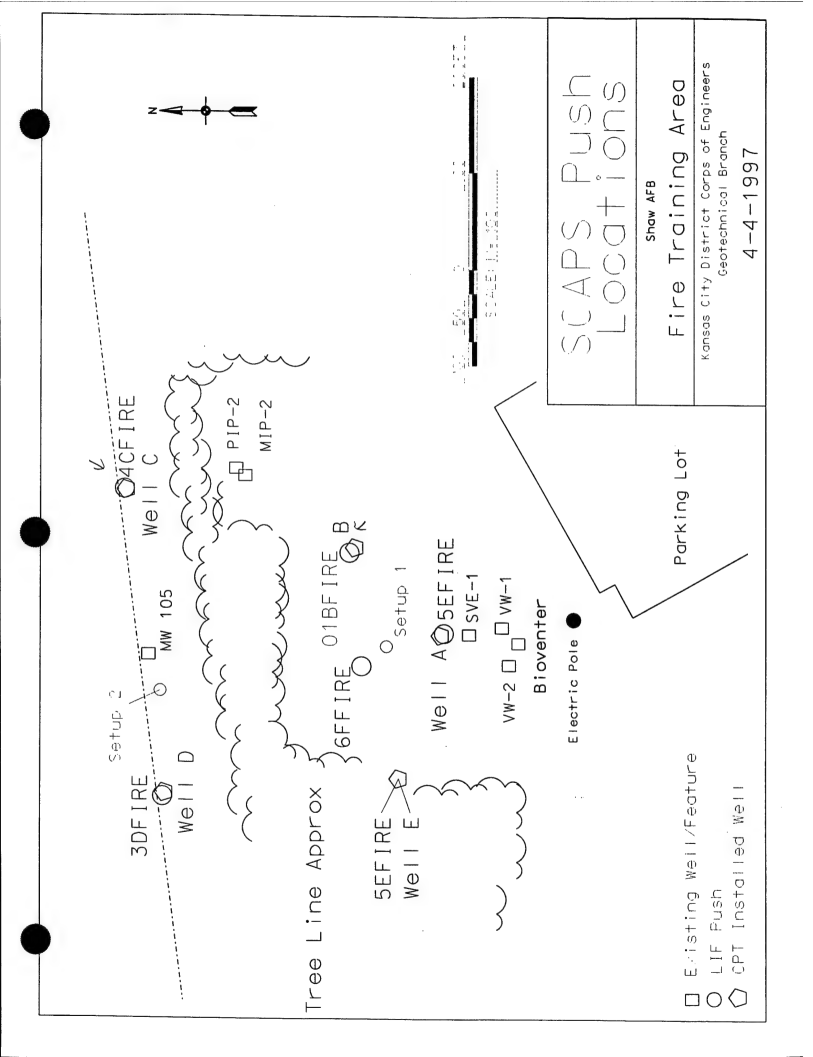
All wells have well boxes which have been cemented in. No bentonite or cement was placed around the well points but the boxes should prevent surface runoff from entering well point.

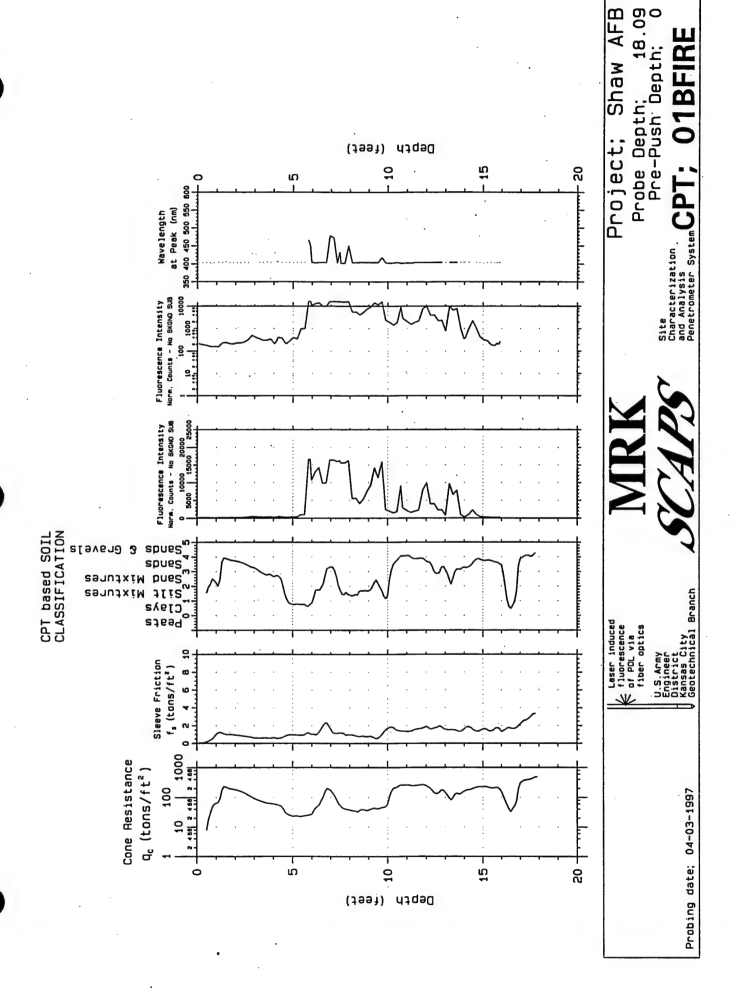
CC:

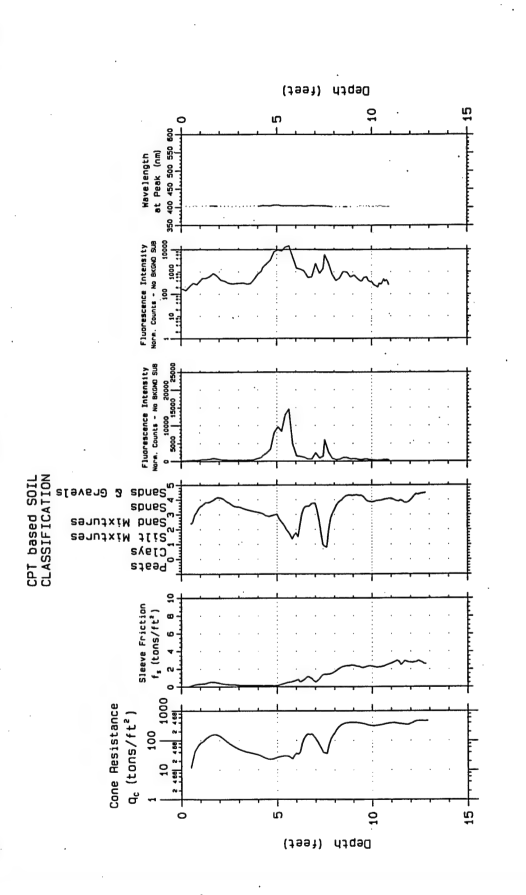
JDC

In addition to the above message sent to Mr. Hansen all well material was schedule 40 flush joint PVC.









Probe Depth; 13 Pre-Push Depth; **2AFIRE** Shaw Project;

13.08 h; 0

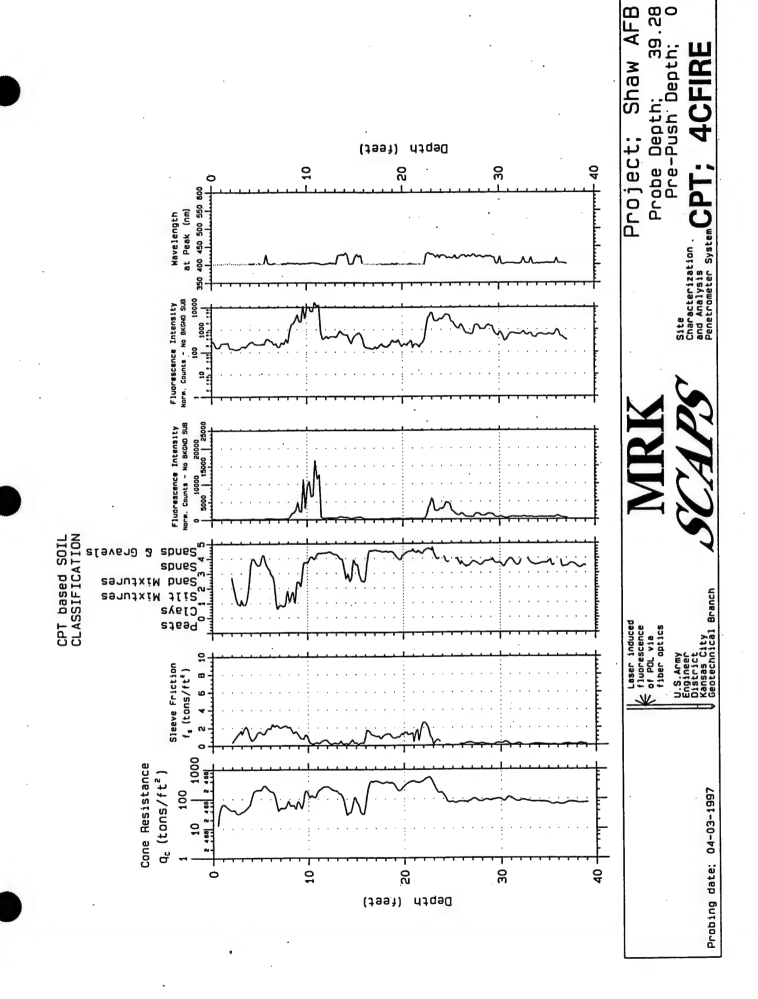
AFB

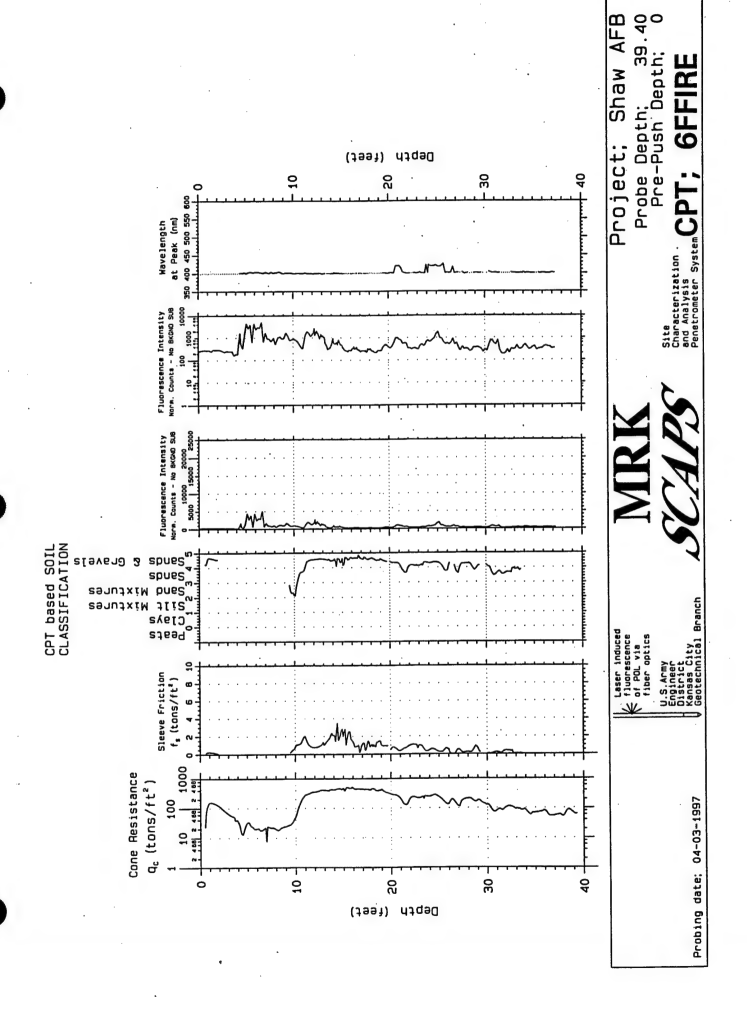
Site Characterization CPT and Analysis Penetrometer System

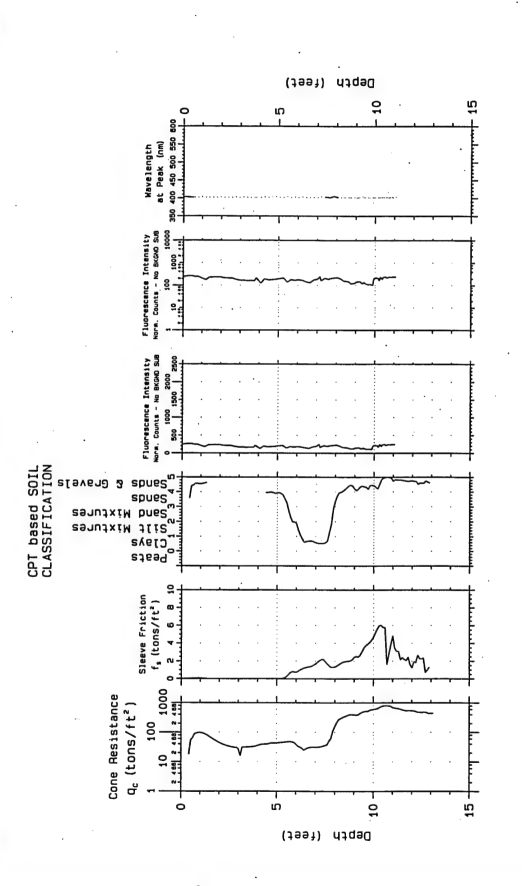
Probing date; 04-03-1997

U.S.Army Engineer District Kansas City Geotechnical Branch

Laser induced fluorescence of POL via fiber optics







13.26 h; 0 AFB Probe Depth; 13 Pre-Push Depth; Shaw Project;

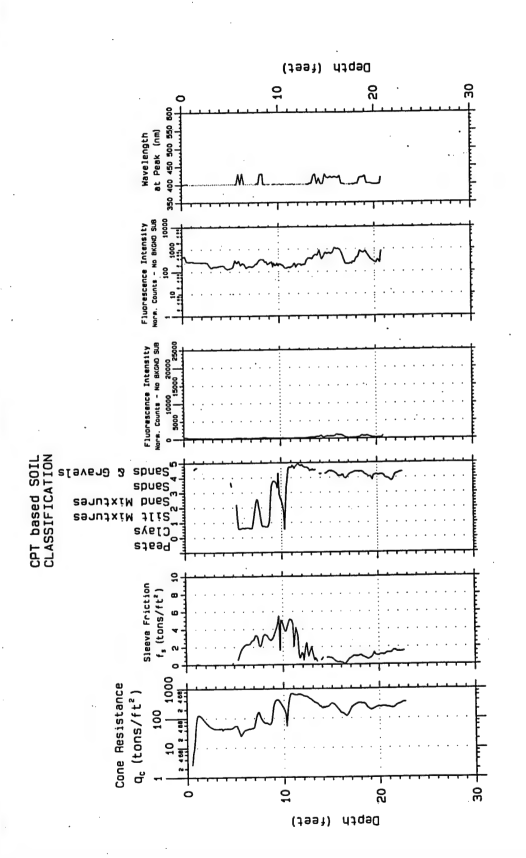
3DFIRE Site
Characterization .

Characterization .

Analysis
Penetrometer System

Probing date; 04-03-1997

U.S.Army Engineer District Kansas City Geotechnical Branch Laser induced fluorescence of POL via fiber optics



22.86 :h: 0 Shaw AFB Probe Depth; 22 Pre-Push Depth; Project;

**SEFIRE** Site Characterization . CPT Penetrometer System

Probing date; 04-03-1997

U.S.Army Engineer District Kansas City Geotechnical Branch Laser induced
fluorescence
of POL via
fiber optics

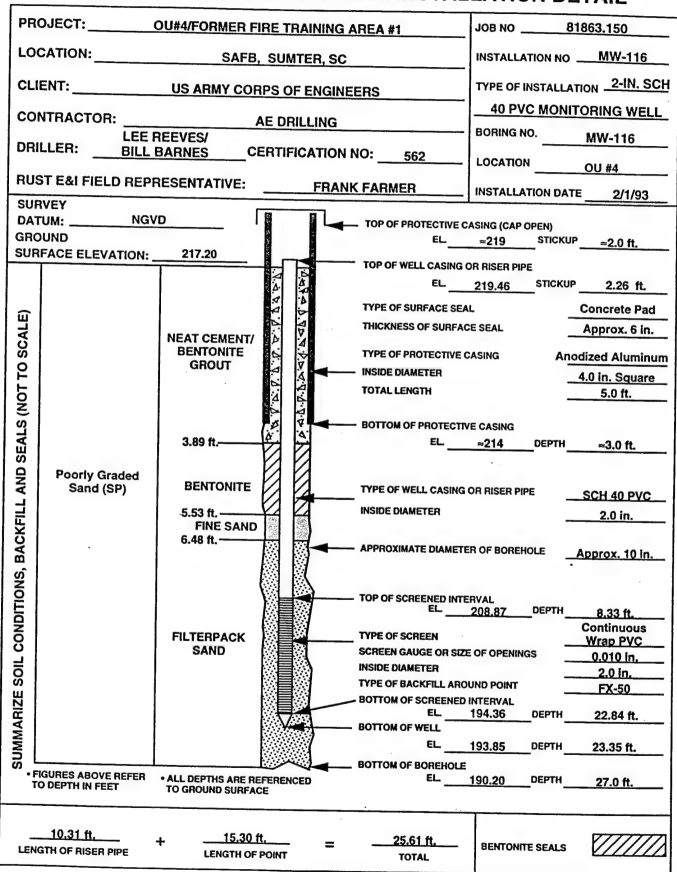


PRO	OJECT:	OU#4/FORMER FIR	E TRAINING A	REA #1	JOB NO81	863.150
						MW-112A
	ENT:	TYPE OF INSTALLA	TION 2-IN. SCH			
1				ENS	40 PVC MONIT	TORING WELL
COI	NTRACTOR:	AE	E DRILLING		BORING NO.	MW-112A
DRI	LLER: B. B.	ARNES CEF	RTIFICATION N	NO: <u>562</u>	LOCATION	
RUS	ST E&I FIELD REPR	ESENTATIVE:	FRANK FA	RMER	INSTALLATION DAT	E 2/4/93
DA1 GRC	RVEY FUM: NGVD DUND RFACE ELEVATION: _	215.78	. V.4 P.V.	TOP OF WELL CASING	≈218 STICKUS OR RISER PIPE	°≈2.0 ft.
			744.7.7.4.7.7.4.7.7.4.4.7.4.4.4.4.4.4.4	TYPE OF SURFACE SEA	AL _	Concrete Pad
SCALE)	1-4	NEAT		THICKNESS OF SURFA	CE SEAL _	Approx. 6 in.
	Interlayered Well Graded and	CEMENT/ BENTONITE	4. A	TYPE OF PROTECTIVE	CASING And	odized Aluminum
(NOT TO	Poorly Graded Sand (SW/SP)	GROUT		- INSIDE DIAMETER	_	4.0 in. Square
TO			4 4	TOTAL LENGTH	_	5.0 ft.
N (S			À	- BOTTOM OF PROTECTI	VE CASING	
SEALS	49.0 ft. ———	60.0 ft.	4		≈213 ВЕРТН	≈3.0 ft.
AND	Well Graded	* BENTONITE		TYPE OF WELL CASING	OR RISER PIPE	SCH 40 PVC 2.0 in.
Ϋ́	Sand (SW)	FINE SAND				
, BACKFILL		73.60 ft.		- APPROXIMATE DIAMET	ER OF BOREHOLE	Approx. 6.5 in.
ONS,		(		- TOP OF SCREENED INT	COVAL	
	78.0 ft.			EL.	136.68 DEPTH_	79.64 ft.
NC				TYPE OF SCREEN		Continuous Slot PVC
S		FILTERPACK		SCREEN GAUGE OR SE	E OF OPENINGS	0.010 in.
OIL	Poorly Graded	SAND	<b>₹</b>	INSIDE DIAMETER	_	2.0 in.
S	Sand (SP)			TYPE OF BACKFILL ARC	_	FX-50
ZIZI		[		-BOTTOM OF SCREENED EL.	126.60 DEPTH	89.18 ft.
SUMMARIZE SOIL CONDIT				BOTTOM OF WELL EL.	126.08 DEPTH	89.70 ft.
S						05.70 16.
• 1	FIGURES ABOVE REFER TO DEPTH IN FEET	ALL DEPTHS ARE REF TO GROUND SURFACE	FERENCED E	BOTTOM OF BOREHOLI	Е <u>125.78</u> DEPTH_	90.00 ft.
LEN	81.51 ft.	+ 10.33 ft,		91.84 ft.	BENTONITE SEALS	



PRO	JECT: OL	J#4/FORMER FIRE T	RAINING AR	EA #1	JOB NO818	63.150
		SAFB, SUM			INSTALLATION NO _	MW-115A
				20	TYPE OF INSTALLAT	ON 2-IN. SCH
	CLIENT: US ARMY CORPS OF ENGINEERS		15	40 PVC MONIT		
CON	TRACTOR:	AE D	RILLING		BORING NO.	MW-115A
DRIL	LER: T. BUR	NETTE CERTI	FICATION NO	D: <u>387</u>	LOCATION	OU #4
RUS	T E&I FIELD REPRE	ESENTATIVE:	FRANK	FARMER	INSTALLATION DATE	2/18/93
SUR				TOP OF PROTECTIVE C	ASING (CAR OREN)	
	JM: NGVI	<u> </u>			≈204 STICKUP	≈2.0 ft.
GRO	UND FACE ELEVATION:	202.19				
SUNI	ACE ELEVATION: _		2	TOP OF WELL CASING		1.97 ft.
	Poorly Graded	À	<u>A</u>	EL	204.16 STICKUP	1.97 π.
	and Well Graded	4	4	TYPE OF SURFACE SEA	\L	Concrete Pad
(E)	Sand (SP/SW)		Ø.	THICKNESS OF SURFA	CE SEAL	Approx. 6. in.
SCALE)	16.0 ft.	NEAT CEMENT/	N P. V	TYPE OF PROTECTIVE	CASING And	odized Aluminum
		GROUT :	<b>4 → →</b>	INSIDE DIAMETER	-	4.0 in. Square
SEALS (NOT TO	Poorly Graded	Ä	4	TOTAL LENGTH		5.0 ft.
Ş	Sand (SP)	. 20	4			
2			D-	BOTTOM OF PROTECT		
AL.	40.6 ft. —	1.4	4.	EL	<u>≈199</u> DEPTH_	≈3.0 ft.
SE,		1.2	\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \			•
AND	Well Graded	47.49 ft.		TYPE OF WELL CASING	OR RISER PIPE	SCH 40 PVC
	Sand (SW)	BENTONITE	4	INSIDE DIAMETER	_	2.0 in.
BACKFILL		54.38 ft.		HODE DIAMETER	_	2.0 111.
X		FINE SAND		- APPROXIMATE DIAME	TER OF BOREHOLE	Approx. 6.5 in.
MA	56.5 ft	56.55 ft.		A Minor	-	
ONS,		l W		TOP OF SCREENED IN	TERVAL 143.70 DEPTH	59.05 ft.
	Clay (CH)	l W			140.70 DEFIN	Continuous
CONDIT		W.	<b>X</b>	TYPE OF SCREEN	_	Slot PVC
ဗ		FILTERPACK (		SCREEN GAUGE OR S	IZE OF OPENINGS	0.010 in.
SOIL	63.0 ft. ———	SAND		INSIDE DIAMETER	OO IND DOINT	2.0 in.
SC	Poorly Graded			TYPE OF BACKFILL AF BOTTOM OF SCREENE		Morie #00N
SUMMARIZE	Sand (SP)			EL_	133.30 DEPTH	68.56 ft.
AR	75.5 ft. ———	1 🛛	<b>V</b>	BOTTOM OF WELL		
M M	Sandy Clay (CL)/			EL_	133.82 DEPTH	69.08 ft.
SC	Clayey Sand (SC)	N. C.		BOTTOM OF BOREHO	LE 113.19 DEPTH	89.0 ft.
	FIGURES ABOVE REFER TO DEPTH IN FEET	• ALL DEPTHS ARE REFE TO GROUND SURFACE	RENCED	<u> </u>	110.13 DEPTH	03.U II.
	60.74 ft.	+ 10.31 ft.	_ =	71.05 ft.	BENTONITE SEALS	· /////
LE	NGTH OF RISER PIPE	LENGTH OF PO	TAIC	TOTAL		<del>_</del>







PR	OJECT:	OU#4/FORMER FIF	RE TRAINING A	REA #1	ЈОВ ИО8	1863.150	
LOCATION: SAFB, SUMTER, SC					INSTALLATION NO	MW-116A	
CLI	ENT:	US ARMY CORE	PS OF ENGINE	ERS	TYPE OF INSTALLATION 2-IN. SCH		
			40 PVC MONITORING WELL				
	LEE F	AREEVES/			BORING NO.	MW-116A	
DRI	LLER: BILL	BARNES CE	RTIFICATION	10. 562	LOCATION		
	ST E&I FIELD REP	RESENTATIVE:	FRANK	FARMER	INSTALLATION DAT		
1	RVEY TUM:NG\	<sub>го</sub> Г		_ TOP OF PROTECTIVE CA	SING (CAR OPEN)		
l l	DUND				≈219 STICKUI	° ≈2.0 ft.	
SUF	RFACE ELEVATION:	217.14		TOP OF WELL CASING O	B BISED DIDE		
			<b>4</b>			2.34 ft.	
	Mainly Poorly		. Ā.				
í í	Graded Sand			TYPE OF SURFACE SEAL	_	Concrete Pad	
Ä	(SP)	NEAT CEMENT/ BENTONITE		THICKNESS OF SURFACE	- SEAL -	6.23 ft.	
SC	49.0 ft.	GROUT		TYPE OF PROTECTIVE C	ASING Ang	dized Aluminium	
2	49.0 ft.		H 3	- INSIDE DIAMETER	_	4.0 in. Square	
SEALS (NOT TO SCALE)			4.4	TOTAL LENGTH	_	5.0 ft.	
2	Well Graded		<b>D</b>	- BOTTOM OF PROTECTIV	ECACING		
ALS	Sand (SW)	72.77 ft.	A. A.		≈214 DEPTH	~30#	
SE,		12.77 10.	7 7 7			-0.0 11.	
AND		BENTONITE					
A	78.0 ft.	79.0 ft. ———		TYPE OF WELL CASING (	OR RISER PIPE _	SCH 40 PVC	
긆		FINE SAND		INSIDE DIAMETER	_	2.0 in.	
BACKFILL		81.65 ft.		49000VIII.			
BA				- APPROXIMATE DIAMETE	R OF BOREHOLE	Approx. 6.5 in.	
(2)							
rions,		· ·		TOP OF SCREENED INTE	RVAL	·	
	Poorly Graded			EL	131,94 DEPTH_	85.20 ft.	
CONDI	Sand (SP)	FILTERPACK SAND		TYPE OF SCREEN		Continuous Wrap PVC	
Ö		J SAILS		SCREEN GAUGE OR SIZE	OF OPENINGS	0.010 in.	
SOIL				INSIDE DIAMETER  TYPE OF BACKFILL AROU	thin power	2.0 in.	
Ш				BOTTOM OF SCREENED	_	FX-50	
RIZ					122.40 DEPTH_	94.74 ft.	
MA				BOTTOM OF WELL	101 90 promi	05.05.4	
SUMMARIZE					121.89 DEPTH_	95.25 ft.	
•	FIGURES ABOVE REFER	ALL DEPTHS ARE RE	FERENCED	BOTTOM OF BOREHOLE	120.14 DEPTH	97.00 ft.	
	TO DEPTH IN FEET	TO GROUND SURFAC	CE				
				· .	1		
	87.26 ft.	+ 10,331	ft	97.59 ft.	BENTONITE SEALS	777777	
LEI	NGTH OF RISER PIPE	LENGTH OF		TOTAL	Sent Office SEALS	K/////	



PR	OJECT:	OU#4/FORMER FIRE	TRAINING A	AREA #1	ЈОВ ИО	11863.150
LO	CATION:	SAFB, SI	UMTER, SC		INSTALLATION NO	MW-117
CLI	ENT:	US ARMY CORPS	OF ENGINE	ERS		TION 2-IN. SCH
co	NTRACTOR:	AE	DRILLING		40 PVC MONI	TORING WELL
	LEE	REEVES/			BORING NO.	MW-117
		BARNES CERT			LOCATION	OU #4
		RESENTATIVE:	FRAN	K FARMER	INSTALLATION DAT	TE <u>2/5/93</u>
DAT	RVEY FUM:NG DUND	VD	1	TOP OF PROTECTIVE C	CASING (CAP OPEN) =216 STICKU	P ≈2.5 ft.
SUR	FACE ELEVATION:	213.89		TOP OF WELL CASING		
		Ž.	1474 b		216.42 STICKU	P2.53 ft.
		2	\$ <del>\Q</del>	TYPE OF SURFACE SEA	AL	Concrete Pad
SCALE)		À	1 N N	THICKNESS OF SURFA	CE SEAL .	Approx. 1.5 ft.
SC.		PROTECTIVE CONCRETE PAD	4	TYPE OF PROTECTIVE	CASING A	nodized Aluminum
5		CONTRICTAD	. ₽ ₩	- INSIDE DIAMETER	-	4.0 in. Square
(NOT	Poorly Graded	4	N.Y.	TOTAL LENGTH		≈3.0 ft.
2	Sand (SP)		4	- BOTTOM OF PROTECTS	WE 04000	
ALS		Δ.	· 4.			≈2.5 ft.
AND SEALS		0.72 ft.				-2.0 14
S		BENTONITE	1 1/2	TYPE OF WELL OLONG		
		2.56 ft.		TYPE OF WELL CASING	OR RISER PIPE	SCH 40 PVC
Ε	3.0 ft	FINE SAND		INSIDE DIAMETER		2.0 in.
BACKFILL	0.010.	3.53 ft.		- APPROXIMATE DIAMET	ER OF BOREHOLF	Anney 10:-
	Olaman On a			•		Approx. 10 in.
ONS,	Clayey Sand (SC)					
		l V		- TOP OF SCREENED INT	ERVAL 207.44 DEPTH	6.45 ft.
SOIL CONDIT	40.74			TYPE OF SCREEN		Continuous
႘	12.5 ft. ———	FILTERPACK SAND		SCREEN GAUGE OR SIZ	Œ OF OPENINGS	Slot PVC 0.010 in.
님		SAND		INSIDE DIAMETER	_	2.0 in.
	Well Graded			TYPE OF BACKFILL ARC		FX-50
SUMMARIZE	Sand (SW)			—BOTTOM OF SCREENED EL.	192.99 DEPTH	20.90 ft.
MA			<b>V</b>	BOTTOM OF WELL		
Σ				EL	192.48 ДЕРТН	21.41 ft.
• F	FIGURES ABOVE REFER TO DEPTH IN FEET	• ALL DEPTHS ARE REFER TO GROUND SURFACE	RENCED	BOTTOM OF BOREHOLE	188.39 DEPTH	25.50 ft.
					<del></del>	
_	8.70 ft.	+ <u>15.24 ft.</u>	_ =	23.94 ft.	DELIFOR THE PARTY OF THE PARTY	777777
LEN	IGTH OF RISER PIPE	LENGTH OF POI		TOTAL	BENTONITE SEALS	V/////



PRO	JECT: O	U#4/FORMER FIRE	TRAINING A	REA #1	JOB NO8	1863.150
LOC	ATION:	SAFB, SU	JMTER, SC		INSTALLATION NO	MW-118
		US ARMY CORPS (			TYPE OF INSTALLA	TION 2-IN. SCH
					40 PVC MONI	TORING WELL
CON	TRACTOR:	AE I	DRILLING		BORING NO.	MW-118
DRIL	LER: T. BUI	RNETTE CERT	IFICATION N	O: <u>387</u>	LOCATION	OU #4
RUS	T E&I FIELD REPR	ESENTATIVE:	FRANK		INSTALLATION DAT	
DAT GRO	VEY UM:NGVE UND FACE ELEVATION: _			TOP OF PROTECTIVE C	≈203 sтіски	P=2.0 ft.
					203.51 стіски	P 2.39 ft.
				TYPE OF SURFACE SEA	•	Concrete Pad
SCALE				THICKNESS OF SURFA	CE SEAL .	Approx. 6 in.
SC/				TYPE OF PROTECTIVE		nodized Aluminum
5			1	INSIDE DIAMETER	•	4.0 in square
(NOT			1 1/1	TOTAL LENGTH	•	Approx. 2.25 ft.
N N		BENTONITE	1	BOTTOM OF PROTECTI	VE CASING	
SEALS	Poorly Graded		1 12	EL	≈200.87 DEPTH	≈0.25 ft.
SE	Sand (SP)		1 M			
AND			1 B	TYPE OF WELL CASING	OR RISER PIPE	SCH 40 PVC
L A			1 1	INSIDE DIAMETER		2.0 in.
FIL			1 12		•	2.0 111.
BACKFILL			1 /4-	APPROXIMATE DIAMET	ER OF BOREHOLE	Approx. 10 in.
, B,	_	0.42 ft.	4 1			•
TIONS,		W.		- TOP OF SCREENED INT	EDVAL	
•		l V		EL	200.27 DEPTH	VIVY III
SOIL CONDI		FILTERPACK		TYPE OF SCREEN		Continuous Wrap PVC
၀		SAND		SCREEN GAUGE OR SE	ZE OF OPENINGS	0.010 in.
٦L		×		INSIDE DIAMETER		2.0 in,
				TYPE OF BACKFILL AR	•	Morie #00N
ZE	8.5 ft.	×.		BOTTOM OF SCREENED EL.	190.76 DEPTH	10.36 ft.
4RI	Clayey Sand (SC)	i i	V	BOTTOM OF WELL		. 3.00 111
MM,	Poorly Graded		<b>\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\</b>	EL.	190.24 DEPTH	10.88 ft.
SUMMARIZE	Sand (SP)	×		BOTTOM OF BOREHOL	E	
•	FIGURES ABOVE REFER TO DEPTH IN FEET	• ALL DEPTHS ARE REFE TO GROUND SURFACE		EL	189.12 DEPTH	12.00 ft.
_	2.96 ft.	10.31 ft	_ =	13.27 ft.	BENTONITE SEALS	
LEI	NGTH OF RISER PIPE	LENGTH OF PO		TOTAL		



PR(	OJECT:		011#4		JOB NO	81961 23N	
LOC	CATION:	SAFB, SO	UTH CAROLIN	Α	INSTALLATION N	o <u>MW-119A</u>	
CLI	ENT:	US ARMY CORPS	S OF ENGINEE	RS		LATION 2-IN. SCH	
	NTRACTOR:				40 PVC MONITORING WELL		
					BORING NO. MW-119A		
DKI	LLER: <u>P. O'D</u>	ONNELL CE	RTIFICATION N	IO: <u>519</u>	LOCATION S	EE SITE PLAN	
RUS	ST E&I FIELD REPR	ESENTATIVE:	B. E.	. CULP	INSTALLATION D		
DAT	RVEY FUM:NGVI DUND RFACE ELEVATION: _			TOP OF PROTECTIVE CASING C	ASING (CAP OPEN) 205.84 ft. STICK		
	Silty Sand (SM)			EL	205.89 ft. STICK	CUP 2.30 ft.	
TTO SCALE)	Well Graded Sand (SW) 20.0————			THICKNESS OF SURFAC — TYPE OF PROTECTIVE O INSIDE DIAMETER TOTAL LENGTH		56.46 ft. nnodized Aluminum 4.0 inch 5.0 feet	
SEALS (NOT	Silty Sand (SM)	Bentonite		BOTTOM OF PROTECTIVE EL.	VE CASING 200.89 ft, DEPTI	H 2.70 ft.	
LL AND	45.0		<b>—</b>	— APPROXIMATE DIAMETI	ER OF BOREHOLE	7 3/4 inch	
IS, BACKFILL	Poorly Graded Sand (SP) 50.5 Silty Sand (SM)	56.46 Fine Sand 57.85		TYPE OF WELL CASING INSIDE DIAMETER	OR RISER PIPE	Schedule 40 PVC 2.0 inch	
DITIONS,	61.0			- TOP OF WELL POINT EL1	122,91 ft. DEPTI	59.92	
RIZE SOIL CONDIT	Clayey Sand (SC)	Filterpack Sand		TYPE OF WELL POINT SCREEN GAUGE OR SIZE INSIDE DIAMETER TYPE OF BACKFILL ARO BOTTOM OF WELL POINT	OUND POINT	Schedule 40 PVC	
SUMMARIZE				EL1 . BOTTOM OF BOREHOLE	133,29 ft. DEPTI	70.30 ft.	
• F	FIGURES ABOVE REFER TO DEPTH IN FEET	ALL DEPTHS ARE REI TO GROUND SURFACE	FERENCED :E		126.59 ft. DEPTH	77.00 ft.	
LEN	62.27 ft.	+ 10.33 f		72.60 ft.	BENTONITE SEAL	s /////	



# SOIL VAPOR EXTRACTION WELL INSTALLATION DETAIL

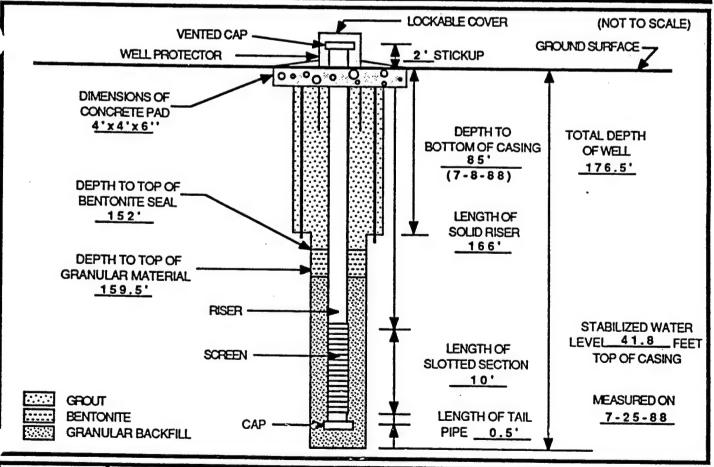
PR	OJECT:	OU#4/FORMER FIRE TR	AINING AREA #1	JOB NO	G-3169.15
LO	CATION:	SAFB, SUM	TER, SC	INSTALLATION NO	SVE-1
CLI	ENT:	US ARMY CORPS OF	ENGINEERS	TYPE OF INSTALL	ATION 4-IN. SCH
1				40 PVC VAPOR E	XTRACTION WELL
CO		AE DRI	LLING	BORING NO.	SVE-1
DRI		BARNES CERTIFI	CATION NO: 562	LOCATION	
RUS	ST E&I FIELD REPF	RESENTATIVE:	FRANK FARMER	INSTALLATION DA	
	RVEY				
	TUM:NG	iVD I		IVE CASING (CAP OPEN) . ≈219 STICKU	IP ~2 #
	RFACE ELEVATION: _	≈216.00 <u> </u>			
			/ B	SING OR RISER PIPE . ≈219 STICKU	1P 2#
í í			TYPE OF SURFAC THICKNESS OF SU	DE40C OCA:	Concrete Pad
SCALE)			THICKNESS OF SC	HPAGE SEAL	6.0 in.
	• '		/ BB	TIVE CASING A	Inodized Aluminum
1			INSIDE DIAMETER		6.0 in. square
101		BENTONITE	TOTAL LENGTH		<u>≈4 ft.</u>
S (P			BOTTOM OF PROT	ECTIVE CASING	
SEALS (NOT TO			EL	≈214 DEPTH	≈2 ft.
SE					
AND			TYPE OF WELL CA	SING OR RISER PIPE	COU 40 DVO
7	Poorly Graded Sand (SP)	2.57 ft.	INSIDE DIAMETER		SCH 40 PVC 4.0 in.
BACKFILL	(3.7)			· .	4.0 111.
ACI			APPROXIMATE DIA	METER OF BOREHOLE	Approx, 10 in.
					•
IONS,			TOP OF WELL POI	ur.	
		FILTERPACK SAND	EL EL	213 DEPTH	2.87 ft.
CONDI			TYPE OF SCREEN		Continuous Wrap PVC
8			SCREEN GAUGE C	R SIZE OF OPENINGS	0.010 in.
등			INSIDE DIAMETER		4.0 in.
ШS			TYPE OF BACKFIL	- AROUND POINT	FX-50
SUMMARIZE SOIL					
MA			BOTTOM OF WELL		
S			EL		8.17 ft.
•	FIGURES ABOVE REFER	• ALL DEPTHS ARE REFEREN	CED BOTTOM OF BORE		9.0 ft.
	TO DEPTH IN FEET	TO GROUND SURFACE			0.0
	5.87 ft.	+ 5.30 ft.	= <u>11.17 ft.</u>	BENTONITE SEALS	
LEI	NGTH OF RISER PIPE	LENGTH OF POINT	TOTAL		

#### TYPE III MONITORING WELL INSTALLATION DIAGRAM

LAW ENVIRONMENTAL, INC. **GOVERNMENT SERVICES DIVISION** KENNESAW, GEORGIA

MW-105D WELL NO. \_ 11-8509-01 JOB NO. \_ 7-13-88 1900 DATE \_ TIME WELL LOCATION SHAW AFB SITE 1. ADJACENT MW-105

GROUND SURFACE ELEVATION ~ 213.16'	BENTONITE TYPE QUIK GEL
TOP OF SCREEN ELEVATION 49.16'	MANUFACTURER NL BAROID DIV.
REFERENCE POINT ELEVATION215.16'	CEMENT TYPE PORTLAND TYPE I MANUFACTURER GIANT CEMENT CO.
TYPE SAND PACK SC#89-M GRADATION GP SAND PACK MANUFACTURER BECKER SAND & GRAVEL	BOREHOLE DIAMETER10" OUTER, 6" INNER
	SCREEN DIAMETER 2" ID SLOT SIZE 0.010
SCREEN MATERIAL PVC-CONTINUOUS WRAP MANUFACTURER JOHNSON	LAW ENVIRONMENTAL, INC. FIELD REPRESENTATIVE MUNGIN-DAVIS
RISER MATERIAL PVC	DRILLING CONTRACTOR LAW ENGINEERING, INC.
MANUFACTURER BRAINARD KILMAN	AMOUNT BENTONITE USED 6 BAGS
RISER DIAMETER 2" ID	
DRILLING TECHNIQUE (above casino) MUD ROTARY	AMOUNT CEMENT USED 29 BAGS
BIT SIZE AND TYPE 9 7/8" TRICONE	AMOUNT SAND USED 10 GALLONS
DRILLING TECHNIQUE (below casing) MUD ROTARY BIT SIZE AND TYPE 5 7/8" TRICONE	STATIC WATER DEPTH (after dev.) 42.20' (TOC) TYPE OF CASING PVC - 6"



A / QC	INSTALLED BY:	R. BANKS	INSTALLATION OBSERVED BY:	QM-D
A / QC	DISCREPANCIES:	STICK UP	2' BEFORE INSTALLATION OF BLADDE	R PUMP

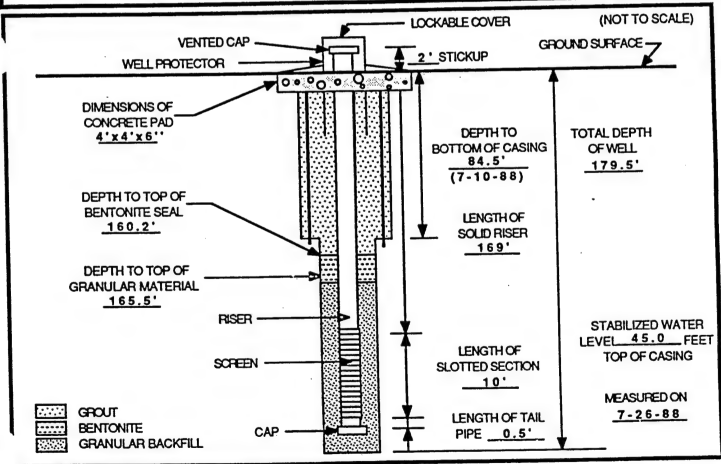
Q

# TYPE III MONITORING WELL INSTALLATION DIAGRAM

LAW ENVIRONMENTAL, INC.
GOVERNMENT SERVICES DIVISION
KENNESAW, GEORGIA

GROUND SURFACE ELE	VATION	~ 217.02'
TOP OF SCREEN ELEVA		50.02'
		219.02'
REFERENCE POINT ELE	VAINN	
TYPE SAND PACK	SC#89-M GR	ADATION GP
SAND PACK MANUFA	CTURER BECKER	SAND & GRAVEL
SCREEN MATERIAL	PVC-CONTIN	UOUS WRAP
MANUFACTURER		1SON
MMMOPACTORET		
RISER MATERIAL	PV	C
• • • • • • • • • • • • • • • • • • • •	DRAINARI	KILMAN
MANUFACTURER _	Bhaillait	RICHIAN
RISER DIAMETER	2"	ID
DRILLING TECHNIQUE	(above casing)	MUD ROTARY
BIT SIZE AND TYPE	9 7/8'	' TRICONE
BIT SIZE AND THE		
DRILLING TECHNIQUE	(below casing) _	MUD ROTARY
		" TRICONE
BIT SIZE AND TYPE	3 770	

QUIK GEL BENTONITE TYPE . NL BAROID DIV. MANUFACTURER -PORTLAND TYPE I CEMENT TYPE \_ GIANT CEMENT CO. MANUFACTURER 10" OUTER, 6" INNER BOREHOLE DIAMETER SCREEN DIAMETER 2" ID SLOT SIZE 0.010 LAW ENVIRONMENTAL, INC FIELD REPRESENTATIVE MUNGIN-DAVIS, ZAUNER DRILLING CONTRACTOR LAW ENGINEERING, INC. AMOUNT BENTONITE USED 8 BAGS AMOUNT CEMENT USED \_\_\_\_\_\_ 26 BAGS 20 GALLONS AMOUNT SAND USED \_\_\_\_\_ STATIC WATER DEPTH (after dev.) 46.14' (TOC) PVC - 6" TYPE OF CASING \_\_\_\_\_

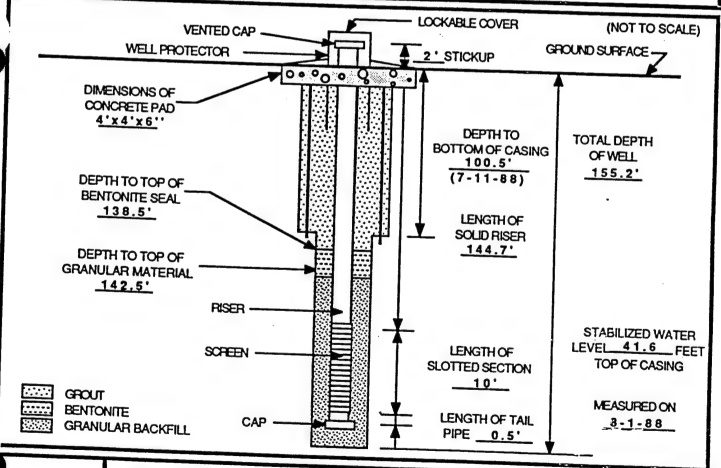


	INSTALLED BY:	C. IVEY	INSTALLATION OBSERVED BY:	QM-D
QA / QC	DISCREPANCIES:		BEFORE INSTALLATION OF BLADDER	PUMP
	DISCHEPANCIES:	0,1.0.0		

# TYPE III MONITORING WELL INSTALLATION DIAGRAM

LAW ENVIRONMENTAL, INC. GOVERNMENT SERVICES DIVISION KENNESAW, GEORGIA 

GROUND SURFACE ELEVATION ~ 215.72'	BENTONITE TYPE QUIK GEL
TOP OF SCREEN ELEVATION73.02'	MANUFACTURER NL BAROID DIV.
REFERENCE POINT ELEVATION217.72'	CEMENTTYPEPORTLAND TYPE
TYPE SAND PACK SC#89-M GRADATION GP SAND PACK MANUFACTURER BECKER SAND & GRAVEL	CORESI DI LE CORESIONE DE LA C
SCREEN MATERIAL PVC-CONTINUOUS WRAP MANUFACTURER JOHNSON	
RISER MATERIAL PVC MANUFACTURER BRAINARD KILMAN	DRILLING CONTRACTOR LAW ENGINEERING, INC.
RISER DIAMETER 2" ID	AMOUNT BENTONITE USED
DRILLING TECHNIQUE (above casing) MUD ROTARY BIT SIZE AND TYPE 9 7/8" TRICONE  DRILLING TECHNIQUE (below casing) MUD ROTARY	AMOUNT CEMENT USED 29 BAGS  AMOUNT SAND USED 15 GALLONS  STATIC WATER DEPTH (after dev.) 44.09' (TOC)
BIT SIZE AND TYPE 5 7/8" TRICONE	TYPE OF CASING PVC - 6" & 10"



QA / QC	INSTALLED BY: C, 1	VEY INCTAL	LATION ODGE	
	DISCREPANCIES:ST		LATION OBSERVED BY:	QM-D
	TIOUTEI AITOIESSI	ICK UP 2. BEFORE	INSTALLATION OF BLA	ADDES BILLIO
				POLIT FOMP

Draw AFB OLLY	Water Levels 5/17/97
Well	DTW (feet)
MW-105	13.54
MW-105D	Couldn't measure
MW-106	17.79
MW-106D	couldn't measure due to deducated pumptubing
MW-107	11.43
MW-111	21.05
MW-112	15.06
MW-112A	19.71
MW-112D	NM (dedicated pump tubing in well-could not get probe past)
MW-113	7.19 yer probe pass
AE11-WM	7.46
MW-114	7.28
MW-115	4.52
MW-115A	7.18
MW-116	15. 28
MW-116A	21.06
MW-117	14.73
MW-118	4.06
TW-101	18.73
MPA	12.92
MP B	14.78
MP C	10.25
MP D	15.18

13-78 500 SHETIS, FILLEN SQUARE
2-381 500 SHETIS THE SACE SSOUNT
2-382 500 SHETIS THE SACE SOUNT
4-382 100 RECYCLED WHITE SSOUNT
4-239 500 RECYCLED WHITE SSOUNT
5-239 500 RECYCLED WHITE SSOU

Job Number: 722450.37020  Location Shaw AFB  Well Identification IYP A	Job Name: AFCEE-RNA by CUMUA PRACE 5/15/97 Measurement Datum TOC 1/2" PVC
Pre-Development Information	Time (Start): 12:30 p
Water Level: 13.5.4	Total Depth of Well: 17.5
Water Characteristics  Color	ature (°C) 22',
Interim Water Characteristics	Well Volume
Gallons Removed 0.5	Well Volume = 0.04 gallon  10 Well Volumes = 0.4 gallons
Temperature (°C) 20,5	
Specific Conductance (45774) 0.580	1 mb/cm
Dissolved Oxygen (mg/L)	<u> </u>
Redox (mV)	
Post-Development Information	Time (Finish): 13:450
Water Level: 13.5	Total Depth of Well: 17.5
Approximate Volume Removed:	.5 gallons
Water Characteristics	Colondon
Color Odor: None Weak	(Ctear Cloudy - Moderate) Strong
Any Films or Immiscible Material	
pH 0,20	Temperature (°C) 21.1
Specific Conductance ( ( )	0.585 m3/om
Dissolved Oxygen (mg/L) (). (2)	
Redox (mV)	
Comments:	

l:\forms\develop.doc

Job Number: 722450,37020  Location Shaw AFB  Well Identification WP B  Job Name: AFCEE-RNA by Liddy   Polity Date: 5/15/9/7  Measurement Datum TOC Va'' PVC
Pre-Development Information Time (Start):
Water Level: did Not register on Solinst Total Depth of Well: did not relastive  Water Characteristics  Color DYMAN Globs in GW Clear Cloudy Odor: None Weak Moderate Strong Any Films or Immiscible Material Yes  PH Temperature (°C) Specific Conductance (µS/cm) Dissolved Oxygen (mg/L) Redox (mV)  Total Depth of Well: did not relastive Solinst was Covered W/ product  Strong full Not measured because foresence of presence of free product
Interim Water Characteristics
Gallons Removed
pH
Temperature (°C)
Specific Conductance(µS/cm)
Dissolved Oxygen (mg/L)
Redox (mV)
Post-Development Information Time (Finish):
Water Level: Total Depth of Well:
Approximate Volume Removed: 2 gallons
Water Characteristics ColorClear Cloudy
Odor: None Weak Moderate Strong  Any Films or Immiscible Material
pHTemperature (°C)
Specific Conductance (µS/cm) Dissolved Oxygen (mg/L)
Redox (mV)
Comments:

i:\forms\develop.doc

Job Number: 722450.37020  Location Shaw AFB  Job Name: AFCEE-RNA by Undy Merrill Date: 5/15/97	
Well Identification MPC Measurement Datum TOC. 72 W PVC	
Pre-Development Information Time (Start): 8:45a	
Water Level: 10.27 Total Depth of Well: 13.62	
Water Characteristics  Color	ゔ
Interim Water Characteristics  3,35 ft WC × 0.01 oct = 0.03	٠.
/ Callana Bamayad	117
Temperature (°C)  Tuck Vikune	-
Temperature (°C)  Specific Conductance (μS/cm)  Dissolved Oxygen (mg/L)  Redox (mV)  NON 0.0335gal = 0,335 gal  Price Victorial  Well did not primp dry at low than product burden to pump and the primp and the primp and the primp and the product burden to pump and the pump and	り 込 -
Post-Development Information Time (Finish): 17 10 6	
Water Level: Total Depth of Well:	
Approximate Volume Removed: 2 gallons	
Water Characteristics  Color Cloudy	
comments: Parameters should be measured during sampling, but we still have background wells to sample 1: Vorms'develop.doc & I den't want to risk cross-contamination by sticking the pickes in the product & water man.	
by sticking the pickes in the product & water now.	

Job Number: 722450.37020  Location Shaw AFB  Job Name: AFCEE-RNA by Lindu Memil Date:	5/15/99
Well Identification MPD Measurement Datum TCC	
Pre-Development Information  Water Level: Total Depth of Wells	
Water Characteristics  Color	0.0537apl
Interim Water Characteristics	1 0
Gallons Removed 1.5 10 Well Vols	=0.537gel
pH	O
1.2	
Redox (mV)	
Redox (mv)	<i>a</i> >-
Post-Development Information Time (Finish):	<u>wp</u>
Water Level: 19.20 Total Depth of Well: 22	2.58
Approximate Volume Removed: 2 gallons	
Water Characteristics  Color Clear Cloudy Odor: None Weak Any Films or Immiscible Material  pH Temperature (°C) 1 K. 3  Specific Conductance (part) 0.026  Dissolved Oxygen (mg/L)  Redox (mV) 25	nelodos
Comments:	

l:\forms\develop.doc

#### GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB Sampling Dates 5/12/97 - 5/22/97

CROIDID II	YATER SAMPLING RECORD - MONITORING WELL
GROUND W	(Identification)
DATE AND SAMPLE CO WEATHER:	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 5/16/97 at 145 a.m./p.m. OLLECTED BY: TH/CM/CH of Parsons ES  Allow A Not 80 F  R WATER DEPTH MEASUREMENT (Describe): TOC /2"   Licon mount
MONITORI	WELL CONDITION:  [ ] LOCKED:  WELL NUMBER (IS - IS NOT) APPARENT  STEEL CASING CONDITION IS:  INNER PVC CASING CONDITION IS:
	WATER DEPTH MEASUREMENT DATUM(IS) IS NOT) APPARENT [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH ISOPROPIS, distilled water Items Cleaned (List): Problem
2 [4]	PRODUCT DEPTH
	WATER DEPTHFT. BELOW DATUM Measured with:
3 [4]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance:  Odor:  Other Comments:
4 13	WELL EVACUATION:  Method:

Monitoring Well No. MPA (Continued) 5 🖂 SAMPLE EXTRACTION METHOD: [] Bailer made of: Verutaltic Other, describe: Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: Measured with Time Temp (°C) (0.05)Cond (MANA) DO (mg/L) Redox (mV) gallons purged ml VOA vials SAMPLE CONTAINERS (material, number, size): 125 ml Volastic A ON-SITE SAMPLE TREATMENT: 8[]8 Containers:\_\_ Method [] Filtration: Containers:\_\_\_ Method Method Containers:\_\_ Preservatives added: [] Containers:\_\_ Method Containers:\_\_\_ Method\_ Containers:\_\_\_ Method Containers:\_\_ Method CONTAINER HANDLING: 9[] Container Sides Labeled [] Container Lids Taped Containers Placed in Ice Chest [] OTHER COMMENTS:\_\_ 10[]

Groundwater Sampling Record

#### GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB
Sampling Dates 5/12/97 - 5/22/97

GROUND W	VATER SAMPLING RECORD - MONITORING WELL VIP 13
	(Identification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [ ] Special Sampling;
DATE AND	TIME OF SAMPLING: 5/16/97 at 330 a.m./p.m.
SAMPLE CO WEATHER:	DLLECTED BY: THICM/CH of Parsons ES hot & sunns 80°F
	R WATER DEPTH MEASUREMENT (Describe): TOC 1/2" ID PVC
	flush mount
	V
MONITORI	NG WELL CONDITION:  [] LOCKED: WUNLOCKED bolted COVER
	WELL NUMBER (IS) IS NOT) APPARENT
	STEEL CASING CONDITION IS: GOOD
	INNER PVC CASING CONDITION IS: GOOD
	WATER DEPTH MEASUREMENT DATUM (IS) IS NOT) APPARENT
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [ ] MONITORING WELL REQUIRED REPAIR (describe):
	[ ] MONITORING WELL REQUIRED RELATIR (describe).
Check-off	LOCADOR DELICATION AND MATER
1[4	EQUIPMENT CLEANED BEFORE USE WITH 1000 CLEANED BEFORE WITH 1000
	Items Cleared (Else).
2 [	PRODUCT DEPTH No 1/2 oil/water interface FT. BELOW DATUM
	Measured with:
	WATER DEPTH NM pencil probe is not FT. BELOW DATUM
	Measured with: Working
/	O COMPANY DEPONE MELL FULL CHATTON (Describe):
3 [U	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: Sultin Grey
	Odor: The strong fuel odor
	Other Comments:
4[1	WELL EVACUATION:  Method: Peristaltic Pump
	Volume Removed: 0.75 Sal
	Observations: Water (slightly) very) cloudy
	Water level (rose fell)- no change) Water odors: Attoma full odor
	Water odors: Attong full gulf Other comments:
	Otter continents.

		Monitori	Froundw	ater San	pling Re	cord (Continue	ed)		
5 12	SAMPLE EX					_(00111111	,		
<b>J</b> [e]		Bailer m	ade of: pe:f escribe:	erista					
6[]	ON-SITE ME	EASUREMEN	NTS:						
	Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV) gallons purged	1.1	3:107 20.1 5.95 0.081 1.0 41 0.35	3:120 19.5 5.92 0.081 0.4 0.4	3:15p 19.6 5:94 0.075 0.4 0.45	3:22p 19.6 6.01 0.079 0.4 -33 0.5	3:26p 19.4 6.01 0.079 0.3 -35 0.55	751 751 751 751 751 750	sured with  3500  3500  3500  3500  3500  4500  51 B  3500
70	SAMPLE CO	2 50 ml 2 105 r 2 350 m	ne <del>sta</del> s	AL ALL	um b Istic	Malae	VOA VI		
8[]	ON-SITE SA	MPLE TREA							
	[]	Filtration:	Meth	od		Contai	ners:		
	[]	Preservatives	added:						
	·			od		Contai	ners: ners: ners:		
9[]	CONTAINE	R HANDLIN	G:						
ر	,	[ ] Conta	iner Sides iner Lids T iners Place	Taped ed in Ice Ch					
10 [4]	OTHER CO	MMENTS:	£	PAY	randl	od 50	mplu	0	

### GROUNDWATER SAMPLING RECORD



Sampling Location Shaw AFB Sampling Dates 5/12/97 - 5/22/97

·	Maria Property Maria Maria	IPC
GROUND WAT	ER SAMPLING RECORD - MONITORING WELL	(Identification)
DATE AND TIMESAMPLE COLL	AMPLING: [X] Regular Sampling; [] Special Sampling; IE OF SAMPLING: 5/16/97 at 10:30 (a.m./p.m. ECTED BY: THOMYCH of Parsons ES	(Identification)
WEATHER: DATUM FOR W	ATER DEPTH MEASUREMENT (Describe): TOC /z"	flish mount
		U
(N)	WELL CONDITION: LOCKED: LOCKED: LOCKED: LY UNLOCKED FELL NUMBER (IS) IS NOT) APPARENT FEEL CASING CONDITION IS:	bolted
/I W 1	NNER PVC CASING CONDITION IS:	Т
Check-off 1 [U E	QUIPMENT CLEANED BEFORE USE WITH USOPADIAL, Items Cleaned (List): Arobis	distilled water
2 M PI	RODUCT DEPTH NM  Measured with: No oil/water interface probe	FT. BELOW DATUM
W	Measured with: (neither Solinst is Working	FT. BELOW DATUM
3 [4 W	ATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance:  Odor:  Other Comments:	
4 [4] W	Method: Peruttalta Pump  Volume Removed: ~ I gollon  Observations: Water (slightly very) cloudy  Water level (rose fell no change)  Water odors:	

Groundwater Sampling Record

Monitoring Well No. (Continued) 5 [1 SAMPLE EXTRACTION METHOD: [] Bailer made of:
Pump, type: Peristaltic Other, describe: Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: Measured with 10: 30 a Time 3900 Temp (°C) 3500 4.99 5.04 рН 5.07 0.060 0.035 Cond (µS/cm) 0.5 DO (mg/L) Redox (mV) 5 callen hillia 0.5 gallons purged 40ml VOA vials SAMPLE CONTAINERS (material, number, size): 50ml class serum olastic bottles 125ml 250ml plastic bottles product Sample ON-SITE SAMPLE TREATMENT: 8[] Containers:\_ Method Filtration: [] Containers:\_\_\_ Method\_ Containers:\_\_ Method Preservatives added: [] Containers:\_ Method Containers:\_\_\_ Method Containers:\_\_\_ Method Containers:\_\_ Method CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped Containers Placed in Ice Chest handled samples OTHER COMMENTS: 10 K

#### GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB Sampling Dates 5/12/97 - 5/22/97

GPOIND W	ATER SAMPLING RECORD - MONITORING WELL
GROOND W	(Identification)
DATE AND SAMPLE CO	R SAMPLING: [X] Regular Sampling; [] Special Sampling:  IIME OF SAMPLING: 5/16/07 at 1/15 a.m./p.m  LLECTED BY: THEMICH of Parsons ES  AMAGIA BOF  R WATER DEPTH MEASUREMENT (Describe): TOC /2" Jush mount
MONITORIN	WELL CONDITION:  [ ] LOCKED:  WELL NUMBER (IS) IS NOT) APPARENT  STEEL CASING CONDITION IS:  INNER PVC CASING CONDITION IS:  WATER DEPTH MEASUREMENT DATUM (IS) IS NOT) APPARENT  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR  [ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH LADRIDGE, distilled water Items Cleaned (List): places
2[4	PRODUCT DEPTH
,	WATER DEPTH NM WOLLIGHT MELL FT. BELOW DATUM  Measured with: 15 110 longer working - See development  Second for previous
3 [1]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: Odor: Other Comments:
4[4	WELL EVACUATION:  Method:  Volume Removed:  Observations:  Water (slightly-very) cloudy  Water level (rose-fell- no change)  Water odors:  Other comments:

		Gr Monitoring	oundwater S	Sampling Re	cord (Continued)		
512	Monitoring Well No. MPD (Continued)  SAMPLE EXTRACTION METHOD:						
<b>3 M</b>	SAME EL	Bailer mad Pump, type Other, desc	le of: e: <u>Purustal</u> cribe:	B; [] COMP	OSITE SAMP		
6 [4]	ON-SITE I	MEASUREMENT	CS:				
ms/cm	Time Temp (°C) pH Cond (#### DO (mg/L) Redox (mV gallons purg	184 5.73 5) 0.042 0 0.7 0.7	100 p   1:03 8.1   18.1 5.64   5.6 0.032   0.04 0.6   0.5 34   36 0.25   0.25	18,1 5,60 1 0.043 0.5 25		Measured with  VSI 3500  VSI 3500  VSI 3500  NSI 51 B  VSI 3500  5 (wllen house)	
714	SAMPLE	CONTAINERS (1 2 50 ml 2 125 m 2 250 m	gass x Lalast	derim t	2. Oller		
8[]	ON-SITE	SAMPLE TREAT	MENT:	J			
	[]	Filtration:	Method Method Method	,	_ Containers:		
	[]	Preservatives a	dded:				
			Method Method Method		Containers Containers		
9[]	CONTAIN	NER HANDLING	:				
10(4)	OTHER C	[ ] Contain	er Sides Labele er Lids Taped ers Placed in Ic		sample	28	

#### GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB Sampling Dates 5/12/97 - 5/22/97

CROLDID II	VATER SAMPLING RECORD - MONITORING WELL
GROUND W	(Identification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [] Special Sampling;
DATE AND	TIME OF SAMPLING: 5/17/97 at 8:45 (a.m./p.m.
SAMPLE CO	DLLECTED BY: TH/CM/CH of Parsons ES
WEATHER:	Clear & breeze 55 t
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): What Measured 14" ID tubing w/strukless steel
	pourt @ 20' bgs
MONITORI	NG WELL CONDITION:
	[] LOCKED:
	WELL NUMBER (IS - IS NOT) APPARENT
	STEEL CASING CONDITION IS:
	INNER PVC CASING CONDITION IS:
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	MONITORING WELL REQUIRED REPAIR (describe):
	/
Check-off	EQUIPMENT CLEANED BEFORE USE WITH 100 propy, distrobed water  Items Cleaned (List): probes
2[1]	PRODUCT DEPTH
-10	Measured with:
	Not Measured
	WATER DEPTHFT. BELOW DATUM
	Measured with:
271	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
3 [[	Appearance: Silty Grey
	Odor: none
	Other Comments:
4[4	WELL EVACUATION: Method: Peristaltic
	1,104:00
	Volume Removed:  Observations: Water(slightly) - very) cloudy
•	Water level (rose -fell) no change)
	Water odors:
	Other comments:
	•

Groundwater Sampling Record Monitoring Well No. MPE (Continued) SAMPLE EXTRACTION METHOD: 5 M [ ] Bailer made of:\_ Pump, type:\_ [ ] Other, describe: Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: Measured with 815a Time <u>35</u>00 Temp (°C) 3500 3500 Cond (#\$%) 0.060 2000 DO (mg/L) Redox (mV) 5 gallon bucket 0.5 gallons purged SAMPLE CONTAINERS (material, number, size): glass serum plastic plastic Naldene bottles ON-SITE SAMPLE TREATMENT: 8[] Containers:\_ Method Filtration: [] Containers:\_\_\_ Method Containers:\_\_ Method Preservatives added: [] Containers:\_\_ Method Containers:\_\_\_ Method Containers:\_\_\_ Method Containers:\_\_\_ Method CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped Containers Placed in Ice Chest OTHER COMMENTS:

GROUND V	VATER SAMPLING RECORD - MONITORING WELL (MP - )	
		dentification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [ ] Special Sampling;	
DATE AND	TIME OF SAMPLING: 5/17 at 1/:04 a.m.p.m.	
	OLLECTED BY: TH/CM/CP of Parsons ES	
WEATHER	sunny ~ 10°	
DATUM FC	OR WATER DEPTH MEASUREMENT (Describe):	
MONITORI	NG WELL CONDITION:	
	[ ] LOCKED: [ ] UNLOCKED	
	WELL NUMBER (IS - IS NOT) APPARENT	
	STEEL CASING CONDITION IS:	
	INNER PVC CASING CONDITION IS:	
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT	
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR	
	[ ] MONITORING WELL REQUIRED REPAIR (describe):	
Check-off		<u> </u>
1[]	EQUIPMENT CLEANED BEFORE USE WITH dedicate	d
- [ ]	Items Cleaned (List):	
•		
	z	
2[]	PRODUCT DEPTH	FT. BELOW DATUM
	Measured with:	
	WATER DEPTH MA	FT. BELOW DATUM
	Measured with:	IT. BELOW DATOM
•	Moderation with.	
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):	
	Appearance:	
	Odor:	
	Other Comments:	
4[]	WELL EVACUATION:  Method: Per: stalt:	
	Volume Removed:	
	Observations: Water (clightly very) cloudy	
•	Water level (rose - fell - no change)	
	Water odors:	
	Other comments:	

Groundwater Sampling Record Monitoring Well No.  $MP - \hat{I}$  (Continued) SAMPLE EXTRACTION METHOD: 5[] [] Bailer made of: peristaltic [ ] Pump, type:\_\_\_\_\_ Other, describe: Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 11:23 Time H:08 11:12 11:16 Temp (°C) 17.0 17.0 19.0 19.1 4.3 4.3 4.3 pН Cond (µS/cm) 20 20 2:1 3.0 DO (mg/L) 2.3 2.6 260 260 Redox (mV) 245 250 235 gallons purged 2 1/2 d SAMPLE CONTAINERS (material, number, size):\_\_\_\_ 7[] 4 VOAS 250-mL ON-SITE SAMPLE TREATMENT: 8[] Containers: Method Filtration: [] Method\_\_\_\_\_ Containers:\_\_\_\_ Containers:\_\_\_ Method Preservatives added: [] Containers: Method Containers:\_\_\_\_ Method Containers: Method\_\_\_ Containers: Method\_\_ 9[] CONTAINER HANDLING: Container Sides Labeled Container Lids Taped

Containers Placed in Ice Chest

OK

OTHER COMMENTS: purged

10[]

GROUND W	ATER SAMPLING RECORD - MONITORING WELL	
01100112 11		(Identification)
REASON FO	OR SAMPLING: [X] Regular Şampling; [] Special Sampling;	
	TIME OF SAMPLING: _//17/67 at 2:15 a.m./p.m.	
SAMPLE CO	DLLECTED BY: TH/CM/CF/ of Parsons ES	
WEATHER.	Sunny ~ 70	
DATIM FO	R WATER DEPTH MEASUREMENT (Describe):	
Ditt Olvi I C		
MONITORI	NG WELL CONDITION:	
	[] LOCKED: [] UNLOCKED	
	WELL NUMBER (IS - IS NOT) APPARENT	
	STEEL CASING CONDITION IS:	
	INNER PVC CASING CONDITION IS:	
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT	
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR	
	MONITORING WELL REQUIRED REPAIR (describe):	
Check-off		
1[]	EQUIPMENT CLEANED BEFORE USE WITH	
	Items Cleaned (List):	
		ET DELOW DATING
2[]	PRODUCT DEPTH	
	Measured with:	
		ET BELOWDATIM
	WATER DEPTH	
	Measured with:	
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):	
2()	Appearance:	
	Odor:	
	Other Comments:	
4[]	WELL EVACUATION:	
	Method:	
	Volume Removed:	
	Observations: Water (slightly - very) cloudy	
	Water level (rose - fell - no change)	
	Water odors:	
	Other comments:	

Groundwater Sampling Record
Monitoring Well No. The Q (Continued) SAMPLE EXTRACTION METHOD: 5[] [ ] Bailer made of:\_ [ ] Pump, type: per: stalt: [ ] Other, describe:\_\_ Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 14:23 14125 Time 17.8 17.9 Temp (°C) 5.1 pН Cond (µS/cm) ひ.ス 0.1 DO (mg/L) · 20 Redox (mV) ~ 1/27=1 gallons purged SAMPLE CONTAINERS (material, number, size):\_ 7[] 254-26 ON-SITE SAMPLE TREATMENT: 8[] Containers:\_\_\_\_ Method\_ Filtration: [] Containers:\_\_\_\_ Method\_\_ Containers:\_\_\_ Method\_\_ Preservatives added: [] Containers: Method Containers: Method Containers:\_\_\_\_\_ Method Containers:\_\_\_\_ Method\_\_\_ CONTAINER HANDLING: 9[] Container Sides Labeled [] Container Lids Taped [] Containers Placed in Ice Chest OTHER COMMENTS:\_\_\_\_\_ 10[]

GROTIND W	ATER SAMPLING RECORD - MONITORING WELL \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	
CICOTID II		(Identification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [ ] Special Sampling;	•
	TIME OF SAMPLING: 5-17-77 at 9:56 am/p.m.	
SAMPLE CO	DLLECTED BY: THCM/CH of Parsons ES	
	Surmy ~ 70	
	R WATER DEPTH MEASUREMENT (Describe):	
Dillomio	(	
MONITORI	NG WELL CONDITION: 52 pant	
	[] LOCKED: [] UNLOCKED	
	WELL NUMBER (IS - IS NOT) APPARENT	
	STEEL CASING CONDITION IS:	
	INNER PVC CASING CONDITION IS:	
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT	
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR	
	MONITORING WELL REQUIRED REPAIR (describe):	
Check-off	~ · · · · · ·	
1[]	EQUIPMENT CLEANED BEFORE USE WITH Dedicated	
	Items Cleaned (List):	
:		ET DELOW DATIN
2[]	PRODUCT DEPTH XIA	F1. BELOW DATUM
	Measured with:	
	WATER DEPTH NA	FT BELOW DATUM
	Measured with:	1 1. DEBO ( D.110)
	Measured with:	
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):	
2[]	Appearance:	
	Odor:	
	Other Comments:	
	Outer Comments.	
4[]	WELL EVACUATION: O ALL	
. ( )	Method: Prishlac	
	Volume Removed:	
	Observations: Water (slightly (very))cloudy	•
	Water level (rose - fell - no change)	
	Water odors: None company	
	Other comments:	

# Groundwater Sampling Record

	Monitoring Well No. <u>Wφ-3</u>	(Continued)					
5[]	SAMPLE EXTRACTION METHOD:						
	[ ] Bailer made of: Porise Lee [ ] Pump, type: [ ] Other, describe:  Sample obtained is [X] GRAB; [ ] COMPO						
[]	ON-SITE MEASUREMENTS:						
		Measured with					
	Time 10:04 10:09 12:1-1 10:16  Temp (°C) 16.6 16.5 16.5						
	Temp (°C)   16.6   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5   16.5						
	Cond (μS/cm) 39 39 39 40						
	Redox (mV) 195 230 350 355 gallons purged 612 20 355						
	ganons purged 12 h 24	Pergy & 200 ML/mm					
7[]	SAMPLE CONTAINERS (material, number, size):						
	4 VoA's						
	2 125 ml/s with Acid						
	I serum with Acid						
	1 20 ML Viol.						
8[]	ON-SITE SAMPLE TREATMENT:						
	[ ] Filtration: Method	Containers:					
	[ ] Filtration: Method	Containers:					
	Method	Containers:					
	[ ] Preservatives added:						
		Containers:					
	Method	Containers:					
		Containers:					
	Method	Containers:					
013	CONTAINER HANDLING:						
9[]	CONTAINER HANDLING.						
	[ ] Container Sides Labeled						
	[ ] Container Lids Taped						
	[ ] Containers Placed in Ice Chest						
10 5 1	OTHER COMMENTS: Pured of.						
10[]	OTHER COMMENTS.						

GPOIND W	WATER SAMPLING RECORD - MONITORING WELL, TMP-4	
GROOND W	(Identification)	
DATE AND SAMPLE CO	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 5/16 at 15:53 a.m./p.m. OLLECTED BY: (HACM/CH) of Parsons ES	
DATUM FO	R WATER DEPTH MEASUREMENT (Describe):	
MONITORIN		
Point	WELL NUMBER (IS - IS NOT) APPARENT STEEL CASING CONDITION IS:	
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR	
Check-off	EQUIPMENT CLEANED BEFORE USE WITH &d: 2 d: 2 d	
2[]	PRODUCT DEPTH	OW DATUM
	WATER DEPTHFT. BELO	OW DATUM
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance:	
4[]	WELL EVACUATION:  Method:	
	REASON FOR DATE AND SAMPLE CONTROL WEATHER DATUM FOR MONITORIA Posicial [1]	STEEL CASING CONDITION IS: INNER PVC CASING CONDITION IS: WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT [] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [] MONITORING WELL REQUIRED REPAIR (describe):  Check-off [] EQUIPMENT CLEANED BEFORE USE WITH

				ater Sam o. TMP		Record (Continued)	)	
5[]	SAMPLE EX	KTRACTION M	1ETHOD	:				
		[ ] Bailer mad [ ] Pump, typ [ ] Other, des  Sample obtaine						
6[]	ON-SITE M	EASUREMEN?	TS:					
	Time	15:58 1	6:04	16:07				Measured with
	Temp (°C)		17.2	17.2				
	pH	4.4	4.3	4.3	1			
	Cond (µS/cm		71	72				
	DO (mg/L)	0.1	0	0				
	Redox (mV)	5	-5	-5				
	gallons purge	d 2/2501		12gal				
7[]	SAMPLE C	ONTAINERS (1	material,	number, siz	ze): c (	(ac.;d)		
8[]	ON-SITE S	AMPLE TREAT	TMENT:					
	[]	Filtration:	Meth	od		Containe	rs:	
	[]	Preservatives a	added:					
			Meth	od		Containe	rs:	
	•		Meth	od		Containe	rs:	
			Meth	odbo		Containe	rs:	
			Meth	odbo		Containe	rs:	

Container Sides Labeled

Container Lids Taped
Containers Placed in Ice Chest

9[]

10[]

CONTAINER HANDLING:

[]

OTHER COMMENTS:\_\_

	GROUND V	vater sampling record - monitoring well TMP - 5	
		a	dentification)
	DATE AND SAMPLE CO	OR SAMPLING: [X] Regular Sampling; [ ] Special Sampling; TIME OF SAMPLING: 5/16 at 14:37 a.m.fp.m. OLLECTED BY: THEMER of Parsons ES	
	DATIM FO	OR WATER DEPTH MEASUREMENT (Describe):	
		AC WITTEN DOI: 111/12/1007-01-12/1007-01-12/1007-01-12/1007-01-12/1007-01-12/1007-01-12/1007-01-12/1007-01-12/	
	MONITORI	NG WELL CONDITION:	
55	Point	[ ] LOCKED: [ ] UNLOCKED  WELL NUMBER (IS - IS NOT) APPARENT  STEEL CASING CONDITION IS:	
	•	INNER PVC CASING CONDITION IS:	
		WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR  [ ] MONITORING WELL REQUIRED REPAIR (describe):	
	Check-off	EQUIPMENT CLEANED BEFORE USE WITH dedicated	
	- ( )	Items Cleaned (List):	
3			
	2[]	PRODUCT DEPTH	FT. BELOW DATUM
		WATER DEPTH	FT. BELOW DATUM
		Measured with:	
	3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: 5/5/1/2 muddy / 60001	
		Odor:	0:3
	4[]	WELL EVACUATION:  Method: peristaltic pump	
		Volume Removed: ~ /OL	
	•	Observations: Water (slightly - very) cloudy Water level (rose - fell - no change)	
		Water odors:	
		Other comments:	

Groundwater Sampling Record

Monitoring Well No. TMP-5 (Continued) SAMPLE EXTRACTION METHOD: 5[] [] Bailer made of:
[] Pump, type:
[] Other, describe: Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 14:42 4:50 14:52 1500 Time 16.2 Temp (°C) 16.3 16 2 4.3 4.5 pН Cond (µS/cm) 47 47 48 0.1 0.1 DO (mg/L) 0.1 0.3 0.4 -30 20 Redox (mV) - 20 gallons purged SAMPLE CONTAINERS (material, number, size):\_\_\_ 7[] 4 VOA (40-mL) 125-mb Plustic (die) 250-mL ON-SITE SAMPLE TREATMENT: 8[] \_\_\_\_\_ Containers:\_\_\_\_\_ Method Filtration: [] Containers: Method\_\_\_\_ Containers:\_\_\_\_ Method Preservatives added: [] Containers: Method\_ Containers:\_\_\_\_ Method\_\_\_ Containers:\_\_\_\_\_ Method Containers: Method\_\_\_ CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped

Containers Placed in Ice Chest

OTHER COMMENTS:\_\_\_\_\_

10[]

GROUND W	ATER SAMPLING RECORD - MONITORING WELL
	(Identification)
DATE AND SAMPLE CO	OR SAMPLING: [X] Regular Sampling; [] Special Sampling;  TIME OF SAMPLING: 5/14/97 at 14'40 a.m./p.m.  OLLECTED BY: THE MCH of Parsons ES - DK4 FB EFA  Partly Cloudy 4 3 10 th Dresse  R WATER DEPTH MEASUREMENT (Describe): 18.70
DATUM FO.	R WATER DEFTH WEASOREWENT (Describe). 1999
) (O) WTODD	IC WELL COMPLETON.
MONITORI	NG WELL CONDITION:  [ ] LOCKED: [M] UNLOCKED
	WELL NUMBER (IS- IS NOT) APPARENT
	STEEL CASING CONDITION IS: Rusty
	INNER PVC CASING CONDITION IS: 4000 WATER DEPTH MEASUREMENT DATUM (15 IS NOT) APPARENT
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	MONITORING WELL REQUIRED REPAIR (describe):
Check-off	
1[]	EQUIPMENT CLEANED BEFORE USE WITH
	Items Cleaned (List):
2[]	PRODUCT DEPTH
~ [ ]	Measured with:
	WATER DEPTH (8.7)FT. BELOW DATUM
	WATER DEPTH 8.7FT. BELOW DATUM  Measured with:
	Mousting Wild.
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
	Appearance: too deep to see
	Other Comments:
4[]	WELL EVACUATION:  Method: Model ES HD DC Parf
	Volume Removed: 35 gal
	Observations: Water (slightly -(very) cloudy
•	Water level (rose - fell - no change)
	Water odors: none
	Other comments: Iron flocularts in early purge water Cleared up at about 10 gal.
	Cleared up at about 10 gm

Groundwater Sampling Record Monitoring Well No. Tw 101 (Continued) SAMPLE EXTRACTION METHOD: 5[] Bailer made of:\_\_\_ My Pump, type: Cole Parmer of a bing part [ ] Other, describe:\_\_\_\_ Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 15/4 1943 1453 1504 Time 21.8 22.6 Temp (°C) 13,1 22,9 6.22 1,23 4.21 6.32 рН 128.7 124.4 125.6 Cond (µS/cm) 132.4 4.1 4,9 4.6 4.0 DO (mg/L) 74.0 71.0 66,4 66.7 Redox (mV) 37.0 gallons purged 35.5 SAMPLE CONTAINERS (material, number, size):\_\_\_\_\_ 7[] ON-SITE SAMPLE TREATMENT: 8[] Containers:\_\_\_\_ Method Filtration: [] Method\_\_\_\_\_ Containers:\_\_\_\_ Containers: Method Preservatives added: [] Containers:\_\_\_\_ Method\_ Containers:\_\_\_\_\_ Method Method\_\_\_\_ Containers:\_\_\_\_ Containers: Method CONTAINER HANDLING: 9[]

- [ ] Container Sides Labeled
- [ ] Container Lids Taped
- [ ] Containers Placed in Ice Chest

10 [ ] OTHER COMMENTS:

Sampling Location Shaw AFB Sampling Dates 5/12/97 - 5/22/97

GROUND W	VATER SAMPLING RECORD - MONITORING WELL
	(Identification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [ ] Special Sampling;
DATE AND	TIME OF SAMPLING: 5/17 at 11:52 (a.m./p.m.
SAMPLE CO	OLLECTED BY: PA/CM/CH) of Parsons ES
WEATHER:	DLLECTED BY: FA/CM/CH of Parsons ES
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): ground surface ( PUC Flush - gro
	, — — — — — — — — — — — — — — — — — — —
MONITORI	NG WELL CONDITION:
	[] LOCKED: [] UNLOCKED
	WELL NUMBER (IS - IS NOT) APPARENT
	STEEL CASING CONDITION IS:
	INNER PVC CASING CONDITION IS:
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	[ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	
[]	EQUIPMENT CLEANED BEFORE USE WITH
	Items Cleaned (List):
	PRODUCT DEPTHFT. BELOW DATUM
2[]	
	WATER DEPTH 5" FT. BELOW DATUM  Measured with: 1402 225212
	WATER DEPTHFT. BELOW DATUM
	WATER DEPTH
	Measured with: type neasure
B[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
	Appearance:
	Odor:
	Other Comments:
[]	WELL EVACUATION:
	Method:
	Volume Removed:
	Observations: Water (slightly - very) cloudy
•	Water level (rose - fell - no change)
	Water odors: Abae
	Other comments:
	Outer comments.

Note: Wall depth = 5 4"

Groundwater Sampling Record Monitoring Well No. Tw-120 (Continued) SAMPLE EXTRACTION METHOD: 5[] Bailer made of: Peristaltic [ ] Other, describe:\_ Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: pumpe ch de-6[] Measured with 12:08 Time 1202 URION Temp (°C) 16.7 16.6 57 Cond (µS/cm) 0.3 DO (mg/L) 0.0 Redox (mV) gallons purged SAMPLE CONTAINERS (material, number, size): 7[] 40-1 VOA (Worsevalive) ON-SITE SAMPLE TREATMENT: 8[] Containers: Method Filtration: [] Containers:\_\_\_ Method Containers:\_\_\_ Method [] Preservatives added: Containers:\_\_ Method Method\_ Containers:\_\_\_\_ Containers:\_\_\_ Method Containers:\_\_ Method CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped [] Containers Placed in Ice Chest OTHER COMMENTS: pursed dry at 12:10pm after jus-10[] Note: Only Filled 4 NOA bottles since nell went dry offer 30 ex purp (somples were muddy)

GROUND W	VATER SAMPLING RECORD - MONITORING WELL
DATE AND SAMPLE CO	(Identification)  OR SAMPLING: [X] Regular Sampling; [] Special Sampling;  TIME OF SAMPLING:5/17 atFr 44 a.m.p.m.  OLLECTED BY: TWCM/CHO of Parsons ES
WEATHER:	R WATER DEPTH MEASUREMENT (Describe): 4.p of 2" PVC (4" ags)
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): 4.0 of 7 POC (9 ags)
MONITORI	NG WELL CONDITION:  [1] LOCKED:  WELL NUMBER (IS - IS NOT) APPARENT  STEEL CASING CONDITION IS:  INNER PVC CASING CONDITION IS:  WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	[ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH
2[]	PRODUCT DEPTHFT. BELOW DATUM Measured with:
	WATER DEPTH 8 1/8 67 ET. BELOW DATUM  Measured with: measuring Tape  FT. BELOW DATUM
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: Odor: Other Comments:
4[]	WELL EVACUATION:  Method:  Volume Removed:  Observations: Water (slightly - very) cloudy
	Water level (rose - fell - no change)  Water odors:  Other comments:
	Note: used tubing from TMP-4 to do  Ha purging  Lell depth = 5 5+

#### Groundwater Sampling Record

Monitoring Well No. TW-(2) (Continued) SAMPLE EXTRACTION METHOD: 5[] [ ] Bailer made of:\_\_ [] Pump, type: peristaltic Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 9:16 4:19 Time 9:07 9:10 4:21 9:07 7.1 17.1 Temp (°C) 17.0 17.1 6.1 6.1 6.0 pН <u> 353</u> Cond (µS/cm) 254 306 342 2.1 4.7 DO (mg/L) smell off Touter Redox (mV) -60 gallons purged - well Ruped thy SAMPLE CONTAINERS (material, number, size): 7[] ON-SITE SAMPLE TREATMENT: 8[] Containers:\_\_\_ Method Filtration: [] Containers:\_\_ Method\_ Containers:\_\_\_ Method\_ Preservatives added: [] Containers:\_ Method Containers:\_\_\_\_ Method Method Containers:\_\_\_ Containers:\_\_\_ Method\_ CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped [] Containers Placed in Ice Chest fter or 2 gallows - west I many OTHER COMMENTS: Well pured Dru 10[]

GROUND W	ATER SAMPLING RECORD - MONITORING WELL Mw 105
OKOOND W	(Identification)
REASON FO	R SAMPLING: [X] Regular Sampling; [ ] Special Sampling;
DATE AND	TIME OF SAMPLING: 5/15/97 at 0800 a.m./p.m.
SAMPLE CO	LLECTED BY: MUNICH OF PRICONES DK & FB & EPA
WEATHER:_	Clear warm
DATUM FOR	R WATER DEPTH MEASUREMENT (Describe): 13.46' from black mark on
casine	
	IC WELT CONDITION:
MONITORIN	NO WELL CONDITION.
	[] LOCKED: MUNLOCKED
· .	WELL'NUMBER(ID- IS NOT) APPARENT
	SILLE CADITO COMPANION IS.
	INNER PVC CASING CONDITION IS: good
	WATER DEPTH MEASUREMENT DATUM (15)- IS NOT) APPARENT
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	[ ] MONITORING WELL REQUIRED REPAIR (describe):
C1 1 CC	
Check-off	EQUIPMENT CLEANED BEFORE USE WITH used new tubing
1[]	EQUIPMENT CLEANED BEFORE USE WITH WSed NEW TRAINS
	Items Cleaned (List):
2[]	PRODUCT DEPTHFT. BELOW DATUM
2[]	Measured with:
	Manager Manage
	WATER DEPTH 13.46 FT. BELOW DATUM
	Measured with: weter probe
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
	Appearance: cloudy (white) with debris in water
	Odor: non observable
	Other Comments:
4[]	WELL EVACUATION:
	Method: periotallic pump
	Volume Removed:
	Observations: Water (slightly - very) cloudy
sec Ne	Water level (rose - Cell- no change)
•	Water odors:
	Other comments: pumped dry at about I gal
	,

# Groundwater Sampling Record

Monitoring Well No. Mw 105 (Continued)

ĺ	Other, complete obta  ASUREME  09/1  19.3  0.29  77.2  0.4	lescribe: ined is [X]	GRAB; [	] COMPO	DSITE SAMPLE	
Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV)	Other, complete obta  ASUREME  09/1  19.3  0.29  77.2  0.4	ined is [X]  NTS:  0816  19.1  5.31  77.3	GRAB; [   0821   19.1   5.31	3825 18.9	OSITE SAMPLE	Measured with
Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV)	Other, complete obta  ASUREME  09/1  19.3  0.29  77.2  0.4	ined is [X]  NTS:  0816  19.1  5.31  77.3	GRAB; [   0821   19.1   5.31	3825 18.9	OSITE SAMPLE	Measured with
Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV)	09/1 19.3 6.29 77.2 0.4	0816 19.1 5.31	0821	3825		Measured with
Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV)	09/1 19.3 6.29 77.2 0.4	0816 19.1 5.31	0821	3825		Measured with
Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV)	0911 19.3 0.29 77.2 0.4	0816 19.1 5.31 77.3	19.1	3825		Measured with
Time Temp (°C) pH Cond (µS/cm) DO (mg/L) Redox (mV)	19.3 6.29 77.2 0.4	19.1 5.31 77.3	19.1	18.9		orion 840
Temp (°C) pH Cond (μS/cm) DO (mg/L) Redox (mV)	19.3 6.29 77.2 0.4	19.1 5.31 77.3	19.1 5.31			
pH Cond (μS/cm) DO (mg/L) Redox (mV)	0.29 77.2 0.4	5.31	5.31	5,31		7004
Cond (µS/cm) DO (mg/L) Redox (mV)	77.2	77.3				
DO (mg/L) Redox (mV)	0.4	0.43		77.1		arion 140
			0.2	0.2		orien 840
gallons purged		100	94.7	.79.0		orion 250 A
	1.5	1.5	1.50 :	1.5		groduated buchet
ON-SITE SAM	PLE TRE	ATMENT:	•			
[ ] F	iltration:	Meth	od		Containers:	
[ ] F	reservative	s added:				
growing on a		Meth	nod nod nod <u></u>	ur have	Containers: Containers: Containers: Containers:	
CONTAINER	HANDLI	NG:		•		
	Conta	ainer Lids 7 ainers Place	Taped'' ed in Ice Ch		high rate	- had to pump
	SAMPLE CON-	SAMPLE CONTAINERS  ON-SITE SAMPLE TRE  [ ] Filtration:  [ ] Preservative  CONTAINER HANDLIN  [ ] Cont.  [ ] Cont.  [ ] Cont.  [ ] Cont.  [ ] Cont.	SAMPLE CONTAINERS (material,  ON-SITE SAMPLE TREATMENT:  [] Filtration: Meth Meth Meth Meth Meth Meth Meth Meth	SAMPLE CONTAINERS (material, number, size of the container sides Labeled [ ] Container Sides Labeled [ ] Container Splaced in Ice Chemical States of the container Splaced in Ice Chemical Splaced in	gallons purged   .5   1.5   1.5   1.5    SAMPLE CONTAINERS (material, number, size):  ON-SITE SAMPLE TREATMENT:  [ ] Filtration: Method	SAMPLE CONTAINERS (material, number, size):  ON-SITE SAMPLE TREATMENT:  [ ] Filtration: Method Containers: M

	WIND SANGE DISCORD MONITORDIS WELL MINI-100
GROUND W	'ATER SAMPLING RECORD - MONITORING WELL (Identification)
DATE AND	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 5/14/9 at 5:00 a.m. fb.m. OLLECTED BY: TH/CM/CH of Parsons ES
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): TOC Z" Q Sticking
MONITORE	NG WELL CONDITION:
WOWTOK	WELL NUMBER (IS IS NOT) APPARENT STEEL CASING CONDITION IS:
	INNER PVC CASING CONDITION IS: 600d
	WATER DEPTH MEASUREMENT DATUM (IS) IS NOT) APPARENT
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH usopropyl, distilled water  Items Cleaned (List): 100 bls
	NA PRIOW DATEM
2[4]	PRODUCT DEPTHFT. BELOW DATUM Measured with:
	WATER DEPTH 17.72 FT. BELOW DATUM Measured with: Solinst With Juri meter
3 M	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: Slightly silty red-brown  Odor: Me Other Comments: —
4[1]	WELL EVACUATION:  Method:  Volume Removed:  Observations:  Water (slightly - very) cloudy  Water evel (rose - fell - no change)  Water odors:  Other comments:

Groundwater Sampling Record Monitoring Well No. MW O6 (Continued) 5 🕅 SAMPLE EXTRACTION METHOD: [ ] Bailer made of: Pump, type:\_ [ ] Other, describe: Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: Measured with Time Temp (°C) Cond (µS/cm) 33400 0.3 DO (mg/L) 1.0 0.5 Redox (mV) -100,8 1.101 --100.2gallons purged SAMPLE CONTAINERS (material, number, size): 4 40ml 7[字 serum bottles (alass) 50ml 125ml plastic Malgene bottles on- site anal ON-SITE SAMPLE TREATMENT: 8[] Containers:\_\_ Method [] Filtration: Containers:\_ Method Containers:\_\_ Method Preservatives added: [] Containers: Method Containers:\_\_ Method\_ Method\_ Containers:\_ Containers:\_\_ Method\_ 9[] CONTAINER HANDLING: Container Sides Labeled [] Container Lids Taped Containers Placed in Ice Chest EPA collected & handled samples

OTHER COMMENTS:

GROUND W	ATER SAMPLING RECORD - MONITORING WELL
01100112 11	(Identification)
DATE AND SAMPLE CO WEATHER:	OR SAMPLING: [X] Regular Sampling; [] Special Sampling;  TIME OF SAMPLING: 5 14 97 at 3:15 a.m.(p.m.)  OLLECTED BY: THICM/CH of Parsons ES  SULAND \$ 80° F  R WATER DEPTH MEASUREMENT (Describe): TOC 2" O PVC STICKUP
————	K WILDER DEL TITLE LOCALIST (CONTROL)
MONITORIN	WELL CONDITION:  [ ] LOCKED:  WELL NUMBER (IS)- IS NOT) APPARENT  STEEL CASING CONDITION IS:  INNER PVC CASING CONDITION IS:  WATER DEPTH MEASUREMENT DATUM (IS)- IS NOT) APPARENT  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR  [ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH LAGROOW , distilled with
2 💜	PRODUCT DEPTH NH FT. BELOW DATUM  Measured with: Solinst Interfall Probe
	WATER DEPTH 11.33FT. BELOW DATUM  Measured with: Soling Interface from
3 [4]	TD = 19.90'  WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance:  Odor:  None
	Other Comments:
4 [4	WELL EVACUATION:  Method:  Volume Removed:  Observations:  Water (slightly -very) cloudy  Water level (rose fell) no change)  Water odors:  Other comments:

Groundwater Sampling Record Monitoring Well No. MW-107 (Continued) SAMPLE EXTRACTION METHOD: [ ] Bailer made of:\_ Pump, type:\_\_ Other, describe: Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE 611 ON-SITE MEASUREMENTS: Measured with 3105 p Time 3:00 P 3:100 Temp (°C) 17.5 17.4 5.47 Ηg 0.026 Cond (<del>µS/em)</del> 0.028 0.027 2.5 2.7 DO (mg/L) Redox (mV) gallons purged 40ml VOA vials SAMPLE CONTAINERS (material, number, size): 4 50 ml alass serum bottles 125 ml plastic Malnere botters ON-SITE SAMPLE TREATMENT: 8[] Containers:\_ Method Filtration: []Containers:\_\_ Method\_ Containers:\_\_ Method Preservatives added: [] Containers: Method Containers:\_\_\_ Method Method\_ Containers:\_ Containers:\_ Method CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped Containers Placed in Ice Chest callected & handled samples OTHER COMMENTS:\_\_

GROUND W	ATER SAMPLING RECORD - MONITORING WELL	MW.	111
			(Identification)
	R SAMPLING: [X] Regular Sampling; [] Special Sampling;		
DATE AND	TIME OF SAMPLING: 5/16/47 at 1017 a.m./p.m.		
SAMPLE CO	DLLECTED BY: PHICM CH-of Parsons ES DK & FB of G	EPA .	
WEATHER:	hazy light wind nice temp.		
DATUM FO	R WATER DEPTH MEASUREMENT (Describe):	arke	
			<u> </u>
MONITORIN	NG WELL CONDITION:		•
	M LOCKED:	KED.	•
	WELL NUMBER (IS- IS NOT) APPARENT		
	STEEL CASING CONDITION IS:		
	INNER PVC CASING CONDITION IS: 9000 ::::		
	WATER DEPTH MEASUREMENT DATUM(19- IS NOT) APP.		Γ
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR		
	[ ] MONITORING WELL REQUIRED REPAIR (describe):		
a			
Check-off	EQUIPMENT CLEANED BEFORE USE WITH new tab	•	
1[]		ing	
	Items Cleaned (List):		
2[]	PRODUCT DEPTH		FT. BELOW DATUM
2[]	Measured with:	-	
	Madada od Mada.		
	WATER DEPTH 21.02		FT. BELOW DATUM
	Measured with: water level meter		
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describ	oe):	
	Appearance:		
•	Odor:		
	Other Comments:		
4[]	WELL EVACUATION:		
	Method: puristaltic pump		
	Volume Removed: 5.052		
•	Observations: Water (slightly - very) cloudy		
	Water level (rose - fell - no change)	) ' (	
	Water odors: none not	cabl	٠
	Other comments:		

# Groundwater Sampling Record

Monitoring Well No. \_\_\_\_\_\_ (Continued)

SAMPLE EXTRACTION METHOD:

5[]	SAMPLE EXT	SAMPLE EXTRACTION METHOD:								
	[ ] Bailer made of:									
	Bailer made of:  Pump, type: Cole Parmer perish the pump  Other, describe:									
	[	] Other, d	lescribe:							
	S	ample obta	ined is [X]	GRAB; [	] COMPO	OSITE SAMPLE				
					•					
6[]	ON-SITE MEA	SUREME 	N15:	• • •	ger i Belgy	· cein ch	\$ 155 MARINE			
	Time	1039	1644	1049	1053		Measured with			
	Temp (°C)	18.9	18.9	19.1	19.0					
	pН	4.796	4.779	4.799						
	Cond (µS/cm)	R.4	19.2	19.1	19.0					
	DO (mg/L)	1.8		8.0	8.0					
	Redox (mV)	208.3	213.2	219						
	gallons purged		1.25	2.0:	2.5					
7[]	SAMPLE CON	ITAINERS	(material,	number, siz	ze):					
, L J			,							
	_									
8[]8	ON-SITE SAM	MPIFTRE	ATMENT:	<b>v</b> erg						
٥١١	ON-SITE SAI	H DD Lim								
	[] F	iltration:	Meth	od	Containers:					
			Meth	od		Containers:				
			Meth	_ Containers:	ontainers:					
	[ ] Preservatives added:									
	. ,									
	•		Meth	od.	Containers:					
			Meth	od		Containers:				
			Meth	od	_ Containers:					
9[]	CONTAINER	HANDLIN	IG:							
	١	] Conta	iner Sides	Labeled						
	•	] Conta	iner Lids T	aped ···		12.47				
	_	] Conta	ainers Place	d in Ice Ch	est					
10.5.1	OTHER COM	MENITO:								
10[]	OTHER COM	MEN 19:								

GROUND W	VATER SAMPLING RECORD - MONITORING WELL MWIIZ
REASON FO DATE AND SAMPLE CO WEATHER:	(Identification)  OR SAMPLING: [X] Regular Sampling; [] Special Sampling;  TIME OF SAMPLING: 5/16/97 at07 46 a.m./p.m.  OLLECTED BY: PHICMICH of Passing ES  Clear at Cool with light wind  R WATER DEPTH MEASUREMENT (Describe): black marker PUC cooling
MONITORI	NG WELL CONDITION:  [ ] LOCKED:  WELL NUMBER (IS- IS NOT) APPARENT  STEEL CASING CONDITION IS:  INNER PVC CASING CONDITION IS:  WATER DEPTH MEASUREMENT DATUM (IS- IS NOT) APPARENT  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR  [ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH new the bing used  Items Cleaned (List):
2[]	PRODUCT DEPTHFT. BELOW DATUM  Measured with:
	WATER DEPTH 15,12' FT. BELOW DATUM  Measured with: water probe
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: Odor: Other Comments:
4[]	WELL EVACUATION:  Method: tubing pump  Volume Removed:  Observations: Water (slightly - very) cloudy not  Water level (rose - fell - no change)  Water odors: Strong manic

# Groundwater Sampling Record

Monitoring Well No. Mw 112 (Continued)

5[]	SAMPLE EX I	RACTION	MEIHOD	•						
	ſ	] Bailer m	nade of:							
	[] Bailer made of:									
	Ĵ	] Other, d	lescribe:	. , ,						
	-		. 1: 52	CD AD. I	1 001/0	OCITE CAMPI	F			
	S	ample obtai	ined is [X]	GRAB; [	COMP	OSITE SAMPL	L			
<i>(</i>	ON-SITE MEA	STIREME	NTS:							
6[]	ON-SITE ME	1001001.00								
	Time	0808	0811	0817	0832	0827	Measured with			
	Temp (°C)	17.9	18.0	18.2	18.2	18.3				
	pH	5.622		5,474		5,403				
	Cond (µS/cm)	108	108	108	108	108				
	DO (mg/L)	0.1	0.0	0,0	0.0	0.0				
	Redox (mV)	-153.9	-1614	-178.2	-183.6	-184.3				
	gallons purged	2.0	2.2	2.5	2.75	3.25				
7[]	SAMPLE CON	VTAINERS	(material,	number, siz	ze):					
	_									
	_									
	-									
0.7.3	ON-SITE SAN	ODIE TRE	ATMENT.							
8[]	ON-SITE SAM	M DE MO.								
	[] · F	iltration:	Meth	od		Containers:_				
	[ ]		Meth	od		Containers:				
			Meth	od		Containers:				
	[ ] F	reservative	es added:							
			Moth	od		Containers:				
			Meth	od		Containers:				
			Meth	od		Containers:				
			Meth	nod		Containers:				
9[]	CONTAINER	HANDLI	NG:							
	•		ainer Sides							
	•		ainer Lids	Taped						
	[	Cont	ainers Place	ed in Ice Ch	est					
	. OTHER COM	(MENITS)								
10[]	OTHER COM	IIMEN I 2:								

GROUND W	WATER SAMPLING RECORD - MONITORING WELL MW 112 A	
		ification)
	OR SAMPLING: [X] Regular, Sampling; [ ] Special Sampling;	•
DATE AND	TIME OF SAMPLING: 5/16/97 at 0746 a.m./p.m.	
	OLLECTED BY: THE OFFICE OF THE SAME TO	
WEATHER:	: Clear of Cool with light Greege	
DATUM FO	OR WATER DEPTH MEASUREMENT (Describe): black mark on Ca	sing
MONITOPI	ING WELL CONDITION:	
MONTOR	[] LOCKED: [] UNLOCKED	
	WELL NUMBER (IS- IS NOT) APPARENT	
	STEEL CASING CONDITION IS: 900 d	
	INNER PVC CASING CONDITION IS: Fair	
	WATER DEPTH MEASUREMENT DATUM(IS)- IS NOT) APPARENT	
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR	
	MONITORING WELL REQUIRED REPAIR (describe):	
Check-off	1	
1[]	EQUIPMENT CLEANED BEFORE USE WITH	
	Items Cleaned (List):	
2[]	PRODUCT DEPTH	T. BELOW DATUM
2[]	Measured with:	
	WATER DEPTH 9.62	T. BELOW DATUM
	Measured with: water probe	
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):	
	Appearance:	
	Odor:	
	Other Comments:	
4 [ ]	WELL EVACUATION:	
4[]	Method: 12 vs + pmp	
	Volume Removed:	
	Observations: Water (slightly - very) cloudy	
•	Water level (rose - fell - no change)	
	Water odors: no ne	
	Other comments:	

# **Groundwater Sampling Record**

Monitoring Well No. Mwいた (Continued)

5[]	SAMPLE EXT	RACTION	METHOD	:				
	] ( 1	Bailer m Pump, ty Other, d	ade of: pe:+u escribe:	Abing pi	<b>ν~</b> Ρ			
	S	ample obtai	ned is [X]	GRAB; [	] COM	POSITE SAM	IPLE	
6[]	ON-SITE MEA	ASUREME	NTS:					
	Time Temp (°C) pH		19,4	18.8				Measured with
	Cond (µS/cm) DO (mg/L) Redox (mV) gallons purged	6.8 105.3	19 6,1 157,7 52	5,6 200,4 54				
7[]	SAMPLE CON	NTAINERS	(material,	number, siz	ze):			
8[]	ON-SITE SAM	IPLE TREA	ATMENT:					
	[] F	iltration:	Meth	od od od		Containe	rs:	
	[] F	reservative	s added:					
			Meth Meth	od od od		Containe Containe	ers: ers:	
9[]	CONTAINER	HANDLIN	G:					
	Ĩ	] Conta	iner Sides I iner Lids T iners Place		est			
10[]	OTHER COM	MENTS:						

CPOIND W	ATER SAMPLING RECORD - MONITORING WELL
GROUND W	(Identification)
DATE AND SAMPLE CO	R SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 5/16/97 at 9:00 (a.m.) p.m.  DLLECTED BY: THCM/CH of Parsons ES  SULLING: 85 F  R WATER DEPTH MEASUREMENT (Describe): TOC 2" () Attuck p
MONITORIN	NG WELL CONDITION:
·	WELL NUMBER (IS)- IS NOT) APPARENT STEEL CASING CONDITION IS: INNER PVC CASING CONDITION IS: WATER DEPTH MEASUREMENT DATUM (IS)- IS NOT) APPARENT [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH 110 propy, distrolod water  Items Cleaned (List): 110 propy, distrolod water
2[4	PRODUCT DEPTH NA FT. BELOW DATUM  Measured with:
	WATER DEPTH 6.5 FT. BELOW DATUM  Measured with:  TD 05' Purge ~ 10 gallons
3 [1]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance:  Odor:  Other Comments:
4 [1]	WELL EVACUATION:  Method:  Volume Removed:  Observations:  Water (slightly - very) cloudy  Water level (rose - fell - no change)  Water odors:  Other comments:

Monitoring Well No. MW-113 (Continued) SAMPLE EXTRACTION METHOD: [ ] Bailer made of:\_ [Y Pump, type: 2 stage [ ] Other, describe:\_ Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 8:40a 1:45a 8:50a Time Temp (°C) pН Cond (10) 0.020 DO (mg/L) Redox (mV) gallons purged 40 ml VOA vials SAMPLE CONTAINERS (material, number, size): 4 40 ml V( plastic Malalne botto senstic Malbere bot ON-SITE SAMPLE TREATMENT: 8[] Containers:\_ Method Filtration: []Method Containers:\_\_\_ Containers:\_\_\_ Method Preservatives added: [] Containers:\_\_ Method\_ Containers:\_\_\_ Method\_ Containers:\_\_\_ Method Containers:\_\_\_ Method CONTAINER HANDLING: 9[] Container Sides Labeled Container Lids Taped Containers Placed in Ice Chest handled samples OTHER COMMENTS:\_

Groundwater Sampling Record

CROIDID WATE	ER SAMPLING RECORD - MONITORING WELL	114
GROUND WATE		(Identification)
SAMPLE COLLE	AMPLING: [X] Regular Sampling; [] Special Sampling; E OF SAMPLING: 5 15 9 at 330 a.m/p.m. ECTED BY: TH/CM/CH of Parsons ES	(Addin Marion)
WEATHER:		Stickey
DATOM FOR W		
NO WEODDIC Y	ARIA COMPITION.	
) WI	VELL CONDITION:  LOCKED:  ELL NUMBER (IS) IS NOT) APPARENT  EEL CASING CONDITION IS:	
	NER PVC CASING CONDITION IS: GCCS	
w. [ ]	ATER DEPTH MEASUREMENT DATUM (IS) IS NOT) APPARENT DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR MONITORING WELL REQUIRED REPAIR (describe):	
l J	MONTORATO WZZZ ZWĘCIAZ ZWIETCZE	
Check-off 1 [ EC	QUIPMENT CLEANED BEFORE USE WITH 15000000000000000000000000000000000000	distilled water
2 [ <b>V</b> PR	CODUCT DEPTH	FT. BELOW DATUM
W	ATER DEPTH 1.27  Measured with: Solinst	FT. BELOW DATUM
3 [4] W.	ATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: Class Odor: More Sutten 199 Other Comments:	S
4 [y] W	Method: 2 Stage Dunch Volume Removed: 10 Gallons Observations: Water (slightly - very) cloudy Water level (rose fell) no change) Water odors: 10 Change	

		<b>Gr</b> een	oundwater Sar g Well No	npling Rec	o <b>rd</b> (Continued)			
5 [4]	SAMPLE EX	XTRACTION M	·					
		[ ] Other, desc	e of:  2 Harl cribe:  d is [X] GRAB;		OSITE SAMPLE			- - -
6 [[]	ON-SITE M	EASUREMENT	S:					
• }	Time	3:00 0 3	10 3:15	3:20		M	leasured with	
	Temp (°C)		7,3 16,7	16.9		Y51	3500	
	pН		4.45 4.9	4.9				
	Cond (µS/cm		100.0 190	0.009			(101	
	DO (mg/L)	0,5. 0	1.5 0.4	10.4		1-1	(YS15B)	
	Redox (mV)	184	73 74	171		1 0	a llon loughest	
	gallons purge	d   4	1 8	1 2		124	alim hudut	İ
7 Y	SAMPLE C	2 105 ml	plastic selection of the plastic	ralaere	office of		in-site and	- - 104815
8[]	ON-SITE SA	AMPLE TREAT			V			
	[]	Filtration:	Method Method Method		Containers: Containers: Containers:			
	[]	Preservatives ac	dded:					
			Method		Containers:			-
			Method		_ Containers:			-
			Method		Containers:			-
			Method		_ Containers:			-
9[]	CONTAINE	ER HANDLING:						
		[] Containe	er Sides Labeled er Lids Taped ers Placed in Ice C					
10 14	OTHER CO	MMENTS: 2	HA ha	ndled	Sample	4		_
* • [Cd					, v			_
								_
								_

GPOLINID W	ATER SAMPLING RECORD - MONITORING WELL MW 115	
GROUND W	ATER SAME BITCHESON MONTH STATES TO THE STATES OF THE STAT	ntification)
REASON FO	R SAMPLING: [X] Regular Sampling; [ ] Special Sampling;	
DATE AND	TIME OF SAMPLING: 5/15/97 at 1215 a.m./p.m.	
SAMPLE CO	Same 115 A	
DATUM FOR	R WATER DEPTH MEASUREMENT (Describe): market on broken	PUC casing
	IG WELL CONDITION:	
MONITORIN	40 WELL COMPINION.	
	[ ] LOCKED: [ ] UNLOCKED WELL NUMBER (19- IS NOT) APPARENT.	
	STEEL CASING CONDITION IS: 9000	
	INNER PVC CASING CONDITION IS! Fair	
	WATER DEPTH MEASUREMENT DATUM ( S- IS NOT) APPARENT	•
	[ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [ ] MONITORING WELL REQUIRED REPAIR (describe):	,
	[ ] MONITORING WELL REQUIRED REPAIR (describe):	
Check-off		
1[]	EQUIPMENT CLEANED BEFORE USE WITH	
	Items Cleaned (List):	
2[]	PRODUCT DEPTH	_FT. BELOW DATUM
	Measured with:	
	WATER DEPTH 4.51	ET BELOW DATIM
	WATER DEPTH	_11. DEEO W DATOM
	Measured with.	
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):	
	Appearance: Clear	
	Odor: strong organic fuel octor Other Comments:	
	Other Comments.	
4[]	WELL EVACUATION:	
	Method: Tubing pump	
	Volume Removed:	
*	Observations: Water (slightly - very) cloudy Water level (rose - fell - no change)	
	Water odors:	
	Other comments:	

# Groundwater Sampling Record Monitoring Well No. MW115 (Continued)

		Sample obta		GRAB; [	] COMP	OSITE SA	MPLE	
]	ON-SITE MEASUREMENTS:					A Third Sale Specials II		
A	Time	1246	1250	1254	1258			Measured with
i	Temp (°C)	17017		17.3	17.3			
	pН	6.099	6.023	5.999	6.003			
	Cond (µS/cm)		2109	211	211			
	DO (mg/L)	0.0	0.0	0.0	0.0			
	Redox (mV)	+190.0	-197.9	-202.5	~ 201.3			
	gallons purge		3.5	4.0.	4.25			
	ON-SITE SA	AMPLE TRE			e):			
l	ON-SITE SA	AMPLE TRE	Meth Meth	odod		_ Contair	ners:	
			Meth Meth Meth	odod		_ Contair	ners:	
	[ ]	Filtration:	Meth Meth Meth es added:	od od od	· · · · · · · · · ·	Contair Contair Contair	ners:	
	[ ]	Filtration:	Meth Meth Meth es added:	od od od	· · · · · ·	Contair Contair Contair Contair	ners:	
	[ ]	Filtration:	Meth Meth Meth es added: Meth Meth	odod		Contair Contair Contair Contair	ners:	
	[ ]	Filtration:	Meth Meth es added: Meth Meth Meth Meth	odododododododod		Contair Contair Contair Contair Contair Contair	ners:	
	[ ]	Filtration: Preservative	Meth Meth Meth es added: Meth Meth Meth	odod		Contair Contair Contair Contair Contair Contair	ners:	
	[]	Filtration:  Preservative  ER HANDLE  [ ] Conta	Meth Meth Meth es added: Meth Meth Meth	odododododod	Special Specia	Contair Contair Contair Contair Contair Contair	ners:	

GROUND W	ATER SAMPLING RECORD - MONITORING WELL 110 115 /7
	(Identification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [ ] Special Sampling;
DATE AND	TIME OF SAMPLING: 5/15/97 at 1215 a.m./p.m.
SAMPLE CO	DLLECTED BY: THICKIET OF PERSON ES DL & FB of EPA
WEATHER:	party Cloudy & warm
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): marked on PVC casing
MONTTORE	NG WELL CONDITION:
MONITORI	
	[ ] LOCKED: [ ] UNLOCKED WELL NUMBER (IS - IS NOT) APPARENT
	STEEL CASING CONDITION IS:
	DILLED CADATO CONDITION IS.
	INNER PVC CASING CONDITION IS:
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	[ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	
1[]	EQUIPMENT CLEANED BEFORE USE WITH
	Items Cleaned (List):
	10110 010110 (210)
2[]	PRODUCT DEPTHFT. BELOW DATUM
	Measured with:
	WATER DEPTH 7.06' FT. BELOW DATUM
	Measured with:
263	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
3[]	Appearance: Clean
	Odor: none
	Other Comments:
	Outer Comments.
4[]	WELL EVACUATION:
4 L J	Method: \$ 12001+440 pump
	Volume Removed:
	Observations: Water (slightly - very) cloudy
•	Water level (rose - fell - no change)
	Water odors:
	Other comments:
	One comment

## Groundwater Sampling Record

	William	Illig well is	0	(Continu	ied)
SAMPLE	EXTRACTION	METHOD	:		
	Bailer r Pump, s Other, o	nade of: type: describe:	white pung		
	Sample obta	ined is [X]	GRAB; []	COMPOSITE S	AMPLE
ON-SITE	MEASUREME			. •	Sanga (1) Ares
Time	1314	1319	1325		Measured
Temp (°C)		19.1	171		
pН	4,821	4.79	4,80		
Cond (µS/c		24	24		
DO (mg/L)		7.4	7.4		
Redox (mV		184.7	196		
gallons pur	ged	<u></u>			
ONLSITE	SAMPLE TRE	ATMENT:			
on-site	SAMPLE TRE	Metho Metho	od	Conta	iners:iners:iners:
		Metho Metho Metho	od	Conta	iners:
[]	Filtration:	Metho Metho Metho es added:	od od	Conta	iners:iners:iners:iners:
[]	Filtration:	Metho Metho Metho es added: Metho Metho	od od od	Conta  Conta  Conta	iners:iners:iners:iners:iners:
[]	Filtration:	Methodes added:  Methodes added:  Methodes Methodes Methodes	od od od od	Conta  Conta  Conta  Conta  Conta  Conta  Conta	iners:in
[]	Filtration:	Methodes added:  Methodes added:  Methodes Methodes Methodes	od od od od	Conta Conta Conta Conta Conta Conta Conta Conta	iners:iners:iners:iners:iners:
[]	Filtration:	Metho Metho es added: Metho Metho Metho	od od od od	Conta  Conta  Conta  Conta  Conta  Conta  Conta	iners:in
[]	Filtration:  Preservative  NER HANDLI  [ ] Conta	Methodes added:  Methodes added:  Methodes Methodetholder  Mothodetholder  Mothodetholder  Methodetholder  Met	od od od od od	Conta Conta Conta Conta Conta Conta Conta Conta	iners:iners:iners:iners:iners:iners:iners:
[]	Filtration:  Preservative  NER HANDLE  [ ] Conta	Methodes added:  Methodes added:  Methodes Methodetholder  Met	odododododododod	Conta Conta Conta Conta Conta	iners:iners:iners:iners:iners:iners:iners:
[]	Filtration:  Preservative  NER HANDLE  [ ] Conta  [ ] Conta  [ ] Conta	Methodes added:  Methodes added:  Methodes Methodetholder  Met	odododododododod	Conta Conta Conta Conta Conta	iners:iners:iners:iners:iners:iners:iners:
[]	Filtration:  Preservative  NER HANDLE  [ ] Conta  [ ] Conta  [ ] Conta	Methodes added:  Methodes added:  Methodes Methodetholder  Met	odododododododod	Conta Conta Conta Conta Conta	iners:iners:iners:iners:iners:iners:iners:

### GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB Sampling Dates 5/12/97 - 5/22/97

GROUND W	VATER SAMPLING RECORD - MONITORING WELL	
GROOND W		entification)
DATE AND SAMPLE CO WEATHER:	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 5/14/97 at 10:00 a.m.(p.m.) OLLECTED BY: TH/CM/CH of Parsons ES  SUMME & WULL  R WATER DEPTH MEASUREMENT (Describe): TOC Z"dag	•
MONITORII	NG WELL CONDITION:    Y LOCKED:   WELL NUMBER (IS) IS NOT) APPARENT   STEEL CASING CONDITION IS:   INNER PVC CASING CONDITION IS:	
	WATER DEPTH MEASUREMENT DATUM (IS (VS NOT) APPARENT  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR  [ ] MONITORING WELL REQUIRED REPAIR (describe):	
Check-off	EQUIPMENT CLEANED BEFORE USE WITH sopropyl, d  Items Cleaned (List): probles	i wales
2 [4]	PRODUCT DEPTH NA  Measured with:	FT. BELOW DATUM
	WATER DEPTH 12.98  Measured with:	FT. BELOW DATUM
3[4	WATER-CONDITION BEFORE WELL EVACUATION (Describe):  Appearance: CLAL  Odor: MONE  Other Comments:	
4[1]	WELL EVACUATION:  Method:  Volume Removed:  Observations:  Water (slightly - very) cloudy  Water level (rose fell) no change)  Water odors:  Other comments:	

	Groundwater, Sampling Record							
/	Monitoring Well No. (Continued)							
5 [4]	SAMPLE EXT	TRACTION	METHOD	):				
	j J		ype: <u>Pe</u> lescribe:			OSITE SAMPLE		
6[4	ON-SITE MEA	ASUREME	NTS:					
	Time	111342	11:40a	11:50a			Measur	red with
	Temp (°C)	21.2	22.3	21.5			Orion	840
0/	pН	NM	4.8	4.6			Orion	250A
mycm->	Cond (µS/cm)	0.05	0.05	0.05			arion	140
	DO (mg/L)	18.5	8.2	8.4			Orion	840
	Redox (mV)	225	234	242			Orión	250A
	gallons purged	1			<u> </u>			
71/	SAMPLE CON	- 50ml	CXUDA O ODO A	tio Valor	Of the land	) ml WOA VU Stles ttles (on-st		156)
8[]	ON-SITE SAM	MPLE TRE	ATMENT:		U			
	[] F	iltration:	Meth	od		Containers:		
			Meth			Containers:		
	•		Meth	od		_ Containers:		
	[ ] F	Preservative	s added:					
			Meth	od		Containers:		
			Meth			_ Containers:		
			Meth			Containers: Containers:		
			Meth	од		_ Containers:		
9[]	CONTAINER	HANDLIN	IG:					
	j	] Conta		aped d in Ice Ch				
10 1/2	OTHER COM	IMENTS:	WA	handle	d samp	لاح		
[0]								

## GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB
Sampling Dates 5/12/97 - 5/22/97

GROUND W	ATER SAMPLING RECORD - MONITORING WELL' MW - 11
GROOND II	(Identification)
DATE AND SAMPLE CO	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING:
WEATHER:	Cloudy - warm  R WATER DEPTH MEASUREMENT (Describe): TOC
DATUMFO	R WATER DEPTH MEASUREMENT (Describe).
•	
MONITORI	NG WELL CONDITION:
	[ ] UNLOCKED.
	WELL NUMBER (IS - IS NOT) APPARENT  STEEL CASING CONDITION IS:  Odd - waver in Haluminum Casing
	STEEL CASING CONDITION IS: 600 C
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	[ ] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	
1[]	EQUIPMENT CLEANED BEFORE USE WITH New to bing
	Items Cleaned (List):
•	
2[]	PRODUCT DEPTHFT. BELOW DATUM
	Measured with:
•	WATER DEPTH 14,66 1 FT. BELOW DATUM
	Measured with: water probe
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
	Appearance: clear
	Odor: Orne
	Other Comments:
4[]	WELL EVACUATION:
	Method: tubing pump
	Volume Removed:
	Observations: Water slightly very) cloudy
	Water level (rose - fell - no change)
	Water odors:
	Other comments:

#### Groundwater Sampling Record

Monitoring Well No. MW117 (Continued) SAMPLE EXTRACTION METHOD: 5[] [ ] Bailer made of:\_ My Pump, type: tubing pump [ ] Other, describe:\_ Sample obtained is [X] GRAB; [ ] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: 6[] Measured with 1635 1138 Time 19.8 Temp (°C) 19.9 pH 4.390 Cond (µS/cm) 61 9.0 9,4 DO (mg/L) 9.5 313,2 3120 ·Redox (mV) gallons purged SAMPLE CONTAINERS (material, number, size):\_\_\_ 7[] ON-SITE SAMPLE TREATMENT: 8[] Containers:\_\_\_\_ Method Filtration: [] Containers: Method\_\_\_ Containers:\_\_\_\_ Method Preservatives added: [] Method .... Containers: Method Containers: \_\_\_\_\_ Containers:\_\_\_\_\_ Method Containers: Method\_\_\_ CONTAINER HANDLING: 9[] Container Sides Labeled
Container Lids Taped [] Containers Placed in Ice Chest [] OTHER COMMENTS:\_\_\_ 10[] 44.3%

# GROUNDWATER SAMPLING RECORD

Sampling Location Shaw AFB
Sampling Dates 5/12/97 - 5/22/97

		MIUIL8	
		TER SAMPLING RECORD - MONITORING WELL : MW 118  (Identification)  (Identification)	ation)
	377 4 7	TR SAMPLING RECORD - MONITORIAN (Identific	attory
C	ROUND WA	Special Sampling;	
		RANGE ING: [X] Regular Sampling; 1 1 Special a.m./p.m.	
I	REASON FOR	SAMPLING: 3/15/97 at 175	
]	DATE AND TI	SAMPLING: [X] Regular Sampling; [] Special Sampling;  ME OF SAMPLING: 5/15/97 at 1409 a.m./p.m.  LECTED BY: What some wind hat not where we are  Clear a hat some wind hat not where we are  WATER DEPTH MEASUREMENT (Describe): black mark an PVC	casing
	SAMPLE COL	LECTED BY: What was hut not where we are PVC  Clear & hot some wind hut not where we are  WATER DEPTH MEASUREMENT (Describe): black mark on PVC	
	WEATHER:	DEPTH MEASUREMENT (Describe)	
	DATUM FOR	WATER DEF III.	
		G WELL CONDITION: [] UNLOCKED.  LOCKED: GO YONGTO APPARENT.	
	MONITORIN	G WELL CONDITION	
	2.2 -	N LOCKED: (R)- IS NOT) APPARENT.	
		WELL CONDITION:  WELL NUMBER (IS- IS NOT) APPARENT  WELL NUMBER (IS- IS NOT) APPARENT  STEEL CASING CONDITION IS:  4 4 5 IS NOT) APPARENT	
		STEEL CASING CONDITION IS: 4 CONDITION APPARENT	
		INNER PVC CASING A SUREMENT DATUM US- IS NOT OF COLLECTOR	
		STEEL CASING CONDITION IS:  INNER PVC CASING CONDITION IS:  WATER DEPTH MEASUREMENT DATUM (IS- IS NOT) APPARENT  WATER DEPTH MEASUREMENT BY SAMPLE COLLECTOR  OF THE CIENCIES CORRECTED BY SAMPLE (describe):	
		WATER DEPTH MEASUREMENT DATUM (IS- IS NOT) THE WATER DATUM (IS- IS NOT) THE W	
		WATER DEPTH MEASUREMENT   WATER DEPTH MEASUREMENT   DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR   DEFICIENCIES CORRECTED REPAIR (describe):	
		EQUIPMENT CLEANED BEFORE USE WITH	
	Check-off	GUEANED BEFORE USE WITH	
	1[]	EQUIPMENT CLEANED BEFORE GSE TO THE SECOND S	
	1 [ ]	Items Cicalica (2007)	FT. BELOW DATUM
			_FT. BELOW DATE
		PRODUCT DEPTH	
,	2[]	PRODUCT DEPTH	_FT. BELOW DATUM
	2[]	Measured Williams	FT. BELOW BITTO
		4.14	
		WATER DEPTH Water probe	
		WATER-CONDITION BEFORE WELL EVACUATION (Describe):	_
		THON REFORE WELL EVACUATION (DESCRIPTION)	
	3[]	WATER-CONDITION BEFORE WELL EVACUATION (CARREST APPEARANCE:	
	2 ( )	Appearance:	
		Other Comments:	
		Other Comments.	
		TO THE OWNER OF THE OWNER	
	4 5 1	WELL EVACUATION:	
	4[]		
		Volume Removed:  Observations: Water (slightly - very) cloudy  Observations: Water (slightly - very) cloudy	
		Water level 11050 1011	
		Water odors:	
		Other comments:	-
ı		Outer comme	
Ŋ.			

Reproduced From Best Available Copy

Groundwater Sampling Record

Monitoring Well No. \_\_\_\_\_\_ (Continued)

5[]	SAMPLE EXT	TRACTION	METHOD	):   -							
	r	] Bailer m	ade of: A								
	Ĭ.	Pump t	vne: Tu	hine Ru	March						
	r I	Other, d	escribe:								
	1	-									
	S	ample obtain	ined is [X]	GRAB;	[] COMP	OSITE SA	MPLE				
6[]	ON-SITE ME	ASUREME.	NTS:	A		· 12 - 15		14:5			
		L	171.2			1446		Measured with			
	Time	1427	1433		1442			1/10454104 //141			
	Temp (°C)	17.8	6.084	17.5	17.5	17.5					
	pH Cond (μS/cm)	6.512			5.883	78					
	DO (mg/L)		78	2.6	2.6	2.5					
	Redox (mV)	162.2	1 4 4		173.6						
	gallons purged		1.5	2.0	2.25		3.0				
	ganons purged	1 I gal	1.3	2.0	, ,,,,		1				
7[]	SAMPLE CO	NTAINERS	(material,	number, si	ze):						
, [ ]	DI MINI DE COI		,								
	_										
	_										
8[]	ON-SITE SAM	MPLE TREA	ATMENT:								
	r., T	iltration:	Math	od		Contai	ners:				
	[ ] F	'iltration:	Meth	od		Contain	Containers:				
			Meth	od		Contai	ners:				
	Method Containers:										
	[] F	[ ] Preservatives added:									
		Method Containers:  Method Containers:									
			Meth	od		_ Contain	ners:				
			Meth	od		Contain	lers.				
			Meth	00		Contail ◆A to/s	ICI 3				
013	CONTAINED	ען ומוא דו	iG·		•	, .,,					
١١٧	CONTAINER	ILANDLIN									
	ſ	] Conta	iner Sides	Labeled							
	•		iner Lids T			1.					
		] Conta	iners Place	d in Ice Ch	est						
		."	5.0								
10[]	OTHER COM	IMENTS:_		•							
				3/4 2 / 4							

## GROUNDWATER SAMPLING RECORD

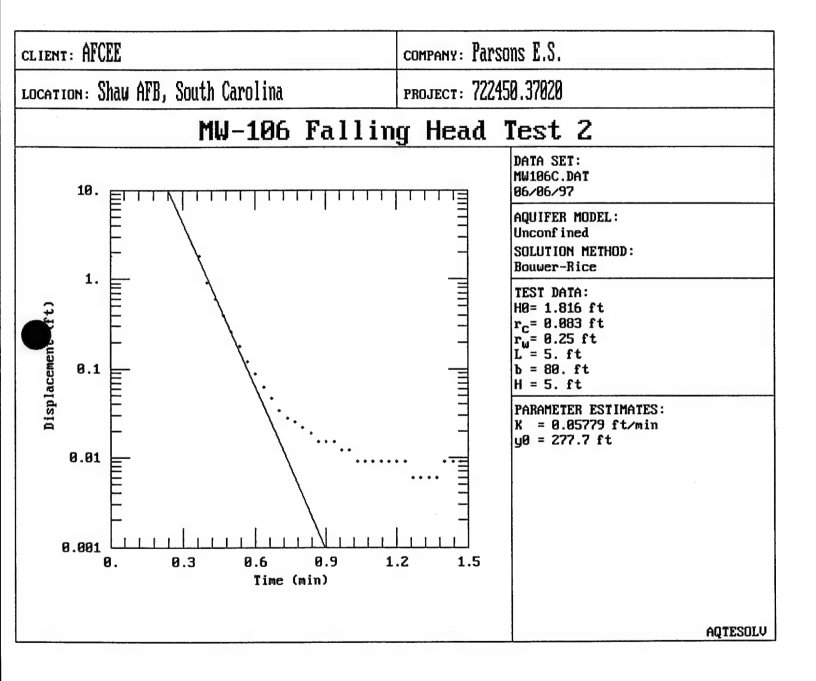
Sampling Location Shaw AFB
Sampling Dates 5/12/97 - 5/22/97

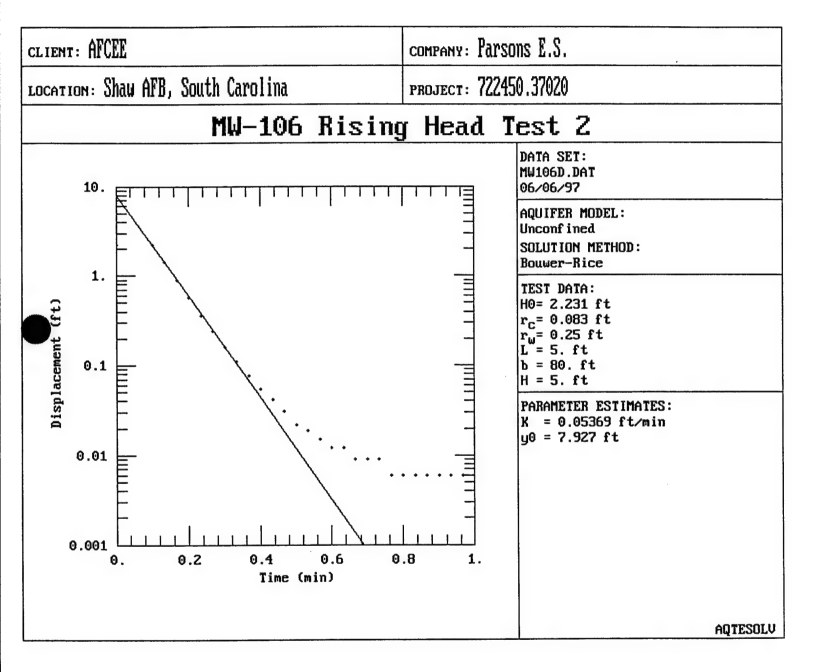
	*
GPOIND V	vater sampling record - monitoring well $\perp \mid \downarrow \langle \varrho A \rangle$
GROOND V	(Identification)
REASON FO	OR SAMPLING: [X] Regular Sampling; [] Special Sampling;
DATE AND	TIME OF SAMPLING: 5/14/97 at 11:30 (a.m./p.m.
SAMPLE CO	DLLECTED BY: TH/CM/CH of Parsons ES
WEATHER:	
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): TOC 2"diam. Stickup
MONITORI	NG WELL CONDITION:
	LOCKED: [ ] UNLOCKED
	WELL NUMBER (IS) IS NOT) APPARENT
	STEEL CASING CONDITION IS: 0000
	INNER PVC CASING CONDITION IS:
	WATER DEPTH MEASUREMENT DATUM (IS (IS NOT) APPARENT NOVEL.  [ ] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
	MONITORING-WELL REQUIRED REPAIR (describe):
Check-off	is a second distinct of with
1 [4	EQUIPMENT CLEANED BEFORE USE WITH ISOMORIS, OUTILLE WITH
	Items Cleaned (List): probles, plush p, Hishing
2 [4	PRODUCT DEPTH NA FT. BELOW DATUM
	Measured with:
	WATER DEPTH 17.79 FT. BELOW DATUM
	WATER DEPTH
	Measured with.
3 M	WATER-CONDITION BEFORE WELL EVACUATION (Describe):
- ( )	Appearance: Ulay
	Odor:
	Other Comments:
4.01	WELL EVACUATION:
4[4	Method: 2 Stage Pump
	Volume Removed: 35 gallons
	Observations: Water (slightly - very) cloudy
	Water level (rose (fell)- no change)
	Water odors:
	Other comments:

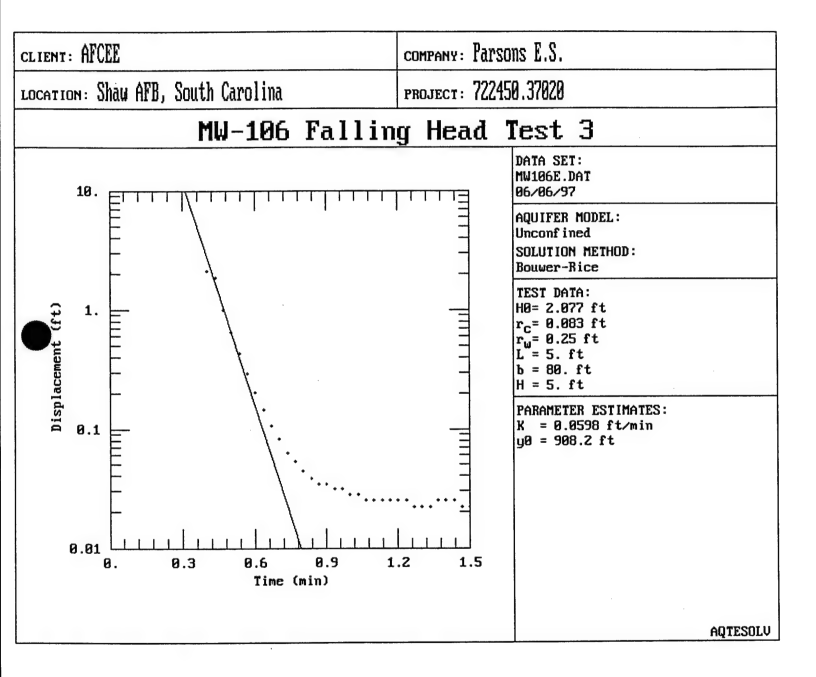
		(	Groundw	vater San	pling Red	cord	
		Monitor	ring Well N	10	A	(Continued)	
54)	SAMPLE EXT	RACTION	METHOD	):			
	i i		ype: lescribe:	GRAB;		OSITE SAME	PLE
614	ON-SITE MEA	SUREME	NTS:				
	Time	10:46a	10:59a	11:07			Measured with
	Temp (°C)	20,0	20.0	20.0			YS1 3500
	рН	5.05	5.08	5.03			1951 3500
mmoh	Cond ( <del>µS/cm)</del>	0.018	0.020	0.190			1/51 3500
/cm	DO (mg/L)	5.2	5.6	5.0			1/51 518
	Redox (mV)	214	255	268			151 3500
	gallons purged	15	20	125			5 gallon bucket
7 M	SAMPLE CON 2 2 2	125 m	Dlast	ic Mal	20):4 40 bottles aene bot Bene bo	tles	viàls -
8[]	ON-SITE SAM	IPLE TRE	ATMENT:				
•	[] F	iltration:	Meth Meth Meth	od		Containers	; ;;
	[ ] P	reservative	s added:				
			Meth	odbo		Containers	:
			Meth	od		_ Containers	
			Meth	od		_ Containers	
			Meth	lod		_ Containers	) <u> </u>
9[]	CONTAINER	HANDLIN	īG:				
/	Ĩ	Conta		Caped d in Ice Ch		onle4	
10 [~]	OTHER COM	MEN 15:_	011	100.00		7	

company: Parsons E.S. CLIENT: AFCEE рвојест: 722450.37020 LOCATION: Shaw AFB, South Carolina MW-106 Falling Head Test 1 DATA SET: MW106A.DAT 06/06/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice TEST DATA: H0= 2.242 ft r<sub>c</sub>= 0.083 ft r<sub>w</sub>= 0.25 ft L = 5. ft 0.1 b = 80. ftH = 5. ftPARAMETER ESTIMATES: K = 0.05498 ft/miny0 = 185. ft0.01 0.001 1.2 1.5 0.3 0.9 Time (min) AQTESOLV

company: Parsons E.S. CLIENT: AFCEE PROJECT: 722450.37020 LOCATION: Shaw AFB, South Carolina MW-106 Rising Head Test 1 DATA SET: MW106B.DAT 06/06/97 10. AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice 1. TEST DATA: H0= 2.021 ft Displacement (ft)  $r_{c} = 0.083 \text{ ft}$ r<sub>w</sub>= 0.25 ft L = 5. ft b = 80. ft0.1 H = 5. ft PARAMETER ESTIMATES: K = 0.05555 ft/miny0 = 283.9 ft0.01 0.001 0.8 1.2 1.6 Time (min) AQTESOLV







company: Parsons E.S. CLIENT: AFCEE LOCATION: Shaw AFB, South Carolina PROJECT: 722450.37020 MW-106 Rising Head Test 3 DATA SET: MW106F.DAT 10. 06/06/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice 1. TEST DATA: H0= 2.503 ft Displacement (ft)  $r_{c} = 0.083 \text{ ft}$ r<sub>w</sub>= 0.25 ft L = 5. ft 0.1 b = 80. ftH = 5. ftPARAMETER ESTIMATES: K = 0.05654 ft/miny0 = 10.27 ft0.01 0.001 0.6 8.0 0.Z 0. Time (min) **AQTESOLV** 

company: Parsons E.S. CLIENT: AFCEE PROJECT: 722450.37020 LOCATION: Shaw AFB, South Carolina MW-112 Falling Head Test 1 DATA SET: MW112A.DAT 06/06/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice TEST DATA: H0= 0.282 ft  $r_c$ = 0.083 ft r<sub>w</sub>= 0.271 ft L"= 15. ft 0.1 b = 80. ftH = 15. ftPARAMETER ESTIMATES: K = 0.03038 ft/miny0 = 8.522 ft0.01 9.001 1.2 1.5 0.6 0.9 0.3 Time (min)

AQTESOLV

company: Parsons E.S. CLIENT: AFCEE рвојест: 722450.37020 LOCATION: Shaw AFB, South Carolina MW-112 Rising Head Test 1 DATA SET: MW112B.DAT 96/96/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice 1. TEST DATA: H0= 1.241 ft  $r_c = 0.083 \text{ ft}$ r<sub>w</sub>= 0.271 ft L = 15. ft Displacemen 0.1 b = 80. ftH = 15. ft PARAMETER ESTIMATES: K = 0.04883 ft/minu0 = 9.81 ft0.01 0.001 8.8 1. 0.2 Time (min) AQTESOLV CLIENT: AFCEE COMPANY: Parsons E.S. LOCATION: Shaw AFB, South Carolina рвојест: 722450.37020 MW-112 Falling Head Test 2 DATA SET: MW112C.DAT 06/06/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice 1. TEST DATA: Displacement (ft) H0= 0.89 ft  $r_c = 0.083 \text{ ft}$ r<sub>w</sub>= 0.271 ft L = 15. ft 0.1 b = 80. ftH = 15. ftPARAMETER ESTIMATES:  $K = 0.03949 \, \text{ft/min}$ y0 = 243.9 ft0.01 0.001 8.3 1.5 0. 1.2 Time (min) AQTESOLV

company: Parsons E.S. CLIENT: AFCEE LOCATION: Shaw AFB, South Carolina рвојест: 722450.37020 MW-112 Rising Head Test 2 DATA SET: MW112D.DAT 10. 06/06/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice TEST DATA: HØ= 1.216 ft r<sub>c</sub>= 0.083 ft r<sub>w</sub>= 0.271 ft L = 15. ft 0.1 b = 80. ftH = 15. ftPARAMETER ESTIMATES: K = 0.05301 ft/miny0 = 37.58 ft0.01 0.001 0.8 0.2 Time (min) AQTESOLV

company: Parsons E.S. CLIENT: AFCEE LOCATION: Shaw AFB, South Carolina PROJECT: 722450.37020 MW-112 Falling Head Test 3 DATA SET: MW112E.DAT 10. 06/06/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice 1. TEST DATA: Displacement (ft) H0= 1.246 ft  $r_c = 0.083 \text{ ft}$ r<sub>ω</sub>= 0.271 ft L = 15. ft b = 80. ft9.1 H = 15. ftPARAMETER ESTIMATES: K = 0.05688 ft/miny0 = 82. ft 0.01 0.001 0.8 0.2 Time (min) AQTESOLV

COMPANY: Parsons E.S. CLIENT: AFCEE LOCATION: Shaw AFB, South Carolina рвојест: 722450.37020 MW-112 Rising Head Test 3 DATA SET: MW112F.DAT 06/86/97 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice TEST DATA: H0= 1.353 ft  $r_c = 0.083 \text{ ft}$ r<sub>w</sub>= 0.271 ft L = 15. ft 0.1 b = 80. ftH = 15. ftPARAMETER ESTIMATES: K = 0.05553 ft/minu0 = 11.66 ft0.01 0.001 0. 0.2 Time (min) AQTESOLV **FAX** 

Ta:

Phone: FAX #



Parsons. ES

303-831-8100 1 303-831-8208

Date:	11 lec 97	
Number o	pages including cover	2



Best Environmental Flight in the Air Force 1994

From:	Parket Koller
	20 CES/CEV
	345 Cullen Street
	Shaw AFB SC 29152-5126
Phone:	(803) 668-5213
FAX#	(803) 868-4383

Routine	Urgent	Review	For Your	Please Reply
		Comment	Information	

SUBJECT:	WELL INFORMATION
Remarks:	Will try to send a more clear
Cl	my late today
	1 85th 1 18

Thanks,

Releaser's Signature



Date:	11 June	7	
Numbe sheet:	r of pages	ncluding cover	2



**FAX** 

То:	Craig Harron or Todd Herrington
	Parsons Engineering
Phone:	303-831-8100, ext1956
FAX#	303-831-8208

From:	Michael	com
	20 CES/0	EV
	345 Culle	Street
	Shaw AF	SC 29152-5126
Phone:	(803) 668	5213, DSN 965-5207
FAX#	(803) 668	4383

SUBJECT: Forwarding well construction info for MW 108 and 112. couldn't get a line the Remarks:

Thanks,

MAN

Releaser's Signature

		-	•	 •		- 5		0 0	- 5	٠. ١	9 6	9
14	CES/CEA	50		Σί	9 <b>2</b> 789	9208	Ιb	:57	<b>حود</b>	st-t	τ-:	90

S0.9

EHAMINIAM MEHAMTABL JOLSWOOTST

u sur			•										rade	Page 8 of 22	
		State Plane Coordinates	Unates	Date	Drilling		Ground	TOC	Stick-Up	TD BGS (ft)	Screen	Screened Interval	2	Bore	83
Location	Well ID	North	East			Aquifer	(ff msl)				€			(E)	
O. H	MAY-70B				Geo-Tech					210	10	200.0	210.0	•	8
OT#2C		775372,5035	2155157.43			S		271.9							
OU#2C		776088.1763	2154649.19			S		288.83							
OU#2C		777058.5003	2155404.63			ST		283.18							
000	-	782128.419	2166707.03			WT2/U	212.92	214.87	1.95	19.7	9	14.7	19.7		7
OL#4	MW-105D	782134.801	2166707.44			BC/L	212.92	215.11	2.19	176.5	9	164.0	174.0		7
0044	MW-106	782452.344	2166601.96			WT2/U	217.63	219.21	1.58	24.3	S.	19.3	24.3		7
OU#4	MW-106D	782448.677	2166595.26			BC/	217.47	218.99	1.52	179.5	9	167.0	177.0		7
OLWA	MW-107	782281,684	2166296.11			WT2/U	213.21	214.79	1.58	19.9	ည	14.9	19.9		7
000	MW-111	782234.008	2103010.20			0514	20.03	-							,
00 12 12	MW-112	782153.317	2166456.63			W120	215.78	218.01	7.73	S	2	0.0	0.07		۷ (
00#4	MW-112A	782153.806	2166465.88			WT2/L	215.78	217.92	2.14	89.7	9.54	79.6	89.2		N (
0U#4	MW-112D	782159.916	2166459.51			BC/L	215.78	217.73	1.95	155.2	£ ;	142.7	152.7		7 (
OU#4	MW-113	782498.827	2166473.27			WT2/U	206.79	208.95	2.16	25	15	5.0	20.0		7
OU#4	MW-113A	782490.479	2166472.45			WT2/L	207.07	209.22	2.15	32	15	20.0	35.0		7
0044	MW-114	782369.261	2166797.62			WT2/U	204.83	207.1	2.27	25	15	5.0	20.0		~
004	MW-115	781886.414	2166877.51	•		WT2/U	201.92	204.19	2.27	25	15	3.0	18.0		<b>N</b>
000	MW-115A	781888.874	2166869.61		•	WTZL	202.19	204.16	1.97	69.08	9.51	59.1	9.89		7
0044	MW-116	781981.943	2166241.08			WT2U	217.20	219.46	2.26	25.6	14.51	8.3	22.8		~
0044	MW-116A	781982.286	2166251.07		-	WT2/L	217.14	219.48	2.34	94.74	9.54	85.2	7.		7
00#4	MW-117	781914.192	2166673.36			WT2/U	213.89	216.42	2.53	21.41	14.45	6.5	20.9		7
00# <b>4</b>	MW-118	781610.669	2166979.11			WT2/U	201.12	203.51	2.39	10.36	9.51	0.0	10.4	•	7
OU#4	PZ-101	781665.342	2166718.03			WT2/U	218.53	220.33	1.8	8	9	20.0	30.0		7
00#4	PZ-102	781686.963	2166705.66			WT2U	218.53	220.19	1.66	28	2	18.0	28.0		8
00#4	PZ-103	781622.498	2166741.98			WT2/U	218.63	220.94	2.31	જ્ઞ	2	25.0	35.0		7
OU#4	M1P1			9/12/96	9/12/96 Geo-Tech	Duplin				32	25	10.0	35.0	9	8
007	M2						9		0	6	Š	000			•
	TW-101	781707.969	2166693.13			WT2	218.08	220.47	2.39	9 9	3 3	20.0	20.0		4 (
CAMES FIFTAS	FT2MW-1	779003.29	2163223.55			WT2/U	226.56	229.59	3.03	76	9.51	14.7	24.2		7 (
OLUS PETAS	FT2MW-10	778780.952	2163123.60			WT2/U.	226.45	229.45	m	25	9.53	13.7	23.3		7
CALINE PITALS	FT2MW-2	778303.598	2163161.72		_•	W12/U	224.09	227.05	2.96	24	9.57	13.4	23.0		7
OLINE FITAS	FT2MW-3	778180.995	2163352.14		•	WT2/U	222.03	225.06	3.03	20	9.54	8.1	17.6		7
OLUG PFTAS	_	778273.665	2163898.15			WT2/U	218.33	221.28	2.95	20	9.6	7.3	16.9		7
COLMING FFETAS	_	778711.639	2163876.21			WT2/U	220.77	223.63	2.88	22	9.54	10.3	19.8		7
CLUS HTAS	_	778496.04	2163399.57			WT2/U	223.00	225.99	2.99	25	9.55	10.4	20.0		7
CLIES FFTAS	_	778721.444	2163531.34			WT2A	223.24	226.36	3.12	\$	9.43	69.6	79.1		7
OLM PFTAS		779000.176	2163701.63		,	WT2/U	222.76	225.64	2.88	23	9.53	11.0	20.5		7
OLM FFA	_	778713.451	2163524.17			WT2/U	223.34	226.38	3.04	24	14.49	8.1	22.6		7

SUBJECT: Parson Es Request for Well defenter

1. Need Top of County + Coordinates for TW-120 30 00-4 TW-121 30 00-9 MW 1602-21 @ 00-9 MW:#4 +#7 @ 57-30 Aloly 16/3

00-4 TW-120 197.64 782021 2167084 TW-121 198.01 781842 2167084

BLDC 1613 MM/613-4 252.29 981790.037 2159779.430 AADS MW1613-7 252.83 781869.579 2159721.743 MADS

Blog 1602 MW 1602-21 250.86 781664.93 2159985.88

212.92 212.92 217.63 218.53 218.63 217.47 215.78 215.78 206.79 204.83 202.19 213.21 225.35 207.07 201.12 215.78 201.92 Prepared for Rust Environmental & Infrastructure -- Shallow Aquifer Monitoring Well Mor. 17, 1993 (top of concrete) 218.08 Concrete Control Monument 93024 MONITORING WELL LOCATIONS 213.21 \* 218.53 213.31 217.75 215.92 206.79 204.83 218.63 213.06 225.55 201.92 218.53 216.10 207.07 217.20 202.35 213.89 201.30 ▲ --- Deep Monitoring Well LEGEND AT SHAW A.F.B. PE DET-CAD 33 ELEV. (top of cosing) 220.47 13 - Piezometer 220.19 220.94 214.87 215.11 219.21 216.42 218.99 227.08 218.01 208.95 207.10 204.19 214.79 217.92 204.16 219.46 217.73 209.22 203.51 CRÔFT

Engineering Company

on R Lavy E. Lani. Et. 384 (BOS) 773-1631 ٠ \$ 2166601.959 2166595.258 2165610.262 2166797.623 2166979.110 2166693.132 2166718.029 2166705.655 2166741.982 2166707.029 2166707.444 2166296.107 2166456.628 2166465.881 2166459.506 2166473.268 2166877.511 2166869,608 2166241.078 2166472.454 EASTING Grade elevation (no concrete) 782234.008 -782153.317 781665.342 781686.963 782448.677 782153.806 782490.479 781707.969 781622.498 782128.419 782134.801 782452.344 782281.684 782159.916 782498.827 782369.261 781886.414 781888.874 781981.943 781610.669 NORTHING Concrete Manument "C-22" (6)
Latitude - 33:58'44" N
Longitude - 80'26'58" W
Elev. = 217.28 .80.27'03" W 80'27'03" W 80'27'00" W 80'27'00" W 80'27'05" W 80'27'13" W 80'27'00" W 80.27.00" W 80.27,00" W 80.27'00" W 80.27.01" W 80'27'01" W 80'27'03" W 80'27'03" W 80'27'03" W 80.26'59" W 80.26'58" W 80'26'58" W 33'58'53" N 80'27'05" W 33'58'53" N 80.27.00" W 80'26'57" W LONGITUDE LAND SURVEYOR 33.58'55" N 33"58"52" N 33.58 49" N 33.58'52" N 32"58"58" N 33.58'57" N 33'58'58" N 33'58'55" N 33'58'52" N 32.58'50" N 33'58'50" N 33'58'55" N 37.58.58 N 33.58'56" N 33.58'55" N 32.58.50 N 37.58'56" N 33.58'58" N 33'58'50" N © PZ-103 ◆ PZ-101 ▼ PZ-101 TW-101 LATITUDE ◆ WW-118 Data on Concrete Control Monuments by Hoffman & Company, Inc. furnished by Shaw A.F.B. (NAD 27) MW-112 MW-112A MW-105D MW-105D MW-106 MW-106D WW-112D MW-113 MW-113A MW-115A MW-116 MW-116A TW-101 PZ-101 MW-107 WW-114 MW-115 MW-118 PZ-102 PZ-103 MW-111 MW-117 # MW-115 4 MW-117 ₩-116A REVISED APRIL 27, 1993 TO SHOW NAD 83 STATE PLANE COORDINATES AND CORRECTED LATITUDE AND LONGITUDE. 600 Concrete Monument "C-21" ( Latitude - 33'58'53" N Longitude - 80'27'08" W Elev. = 222.85 MW-1050 ▲ WW-105 MW-112DA WW-112A 1" = 200◆ WW-107 . 00 10 ♣ MW-114 MW-106 SCALE: MW-113 40 MW-113A 200 100

## APPENDIX B

SOIL, SEDIMENT, GROUNDWATER, AND SURFACE WATER ANALYTICAL RESULTS COLLECTED AS PART OF THIS TS.

# MANTECH POST OFFICE BOX 1198 ADA, OK 74820

	FRX PHONE: 405-436-8501	DERIFICATION PHONE: 405-436-8660
	DATE: 6/19/97	#PAGES (INCLUDING COVER SHEET) 2
	To: Craig Ha	kran
)	PHONE:	FAX: 303-83/-8208
	FROM: Sharen PHONE: 405-4	Hightower 36-8664 De 436-8682
	COMMENTS:	

405-436-8564

Dr. Don Kampbell

National Risk Management Research Laboratory Subsurface Protection & Remediation Division

U.S. Environmental Protection Agency

P.O. Box 1198

Ada, OK 74820

compound islandification

THRU: S.A. Vandegrift 🗸

Dear Dr. Kampbell:

This report contains the results of my GC/MSD analysis of the following samples from Shaw AFB under Service Request #SF-3-266:

- Floating product samples
  - 1. OU-4 MPB
  - 2. OU-4 MPC
  - 3. MW1613-12

for compound identification by mass spectral library searching and chromatography of known standards. TIC chromatograms are also included.

- II. Methylene chloride core extracts
  - 1. SED-1
  - 2. SED-2
  - 3. SED-3
  - 4. SED-4
  - 5. SED-5

for quantitative determination of benzene, toluene, ethylbenzene, xylene isomers, trimethylbenzene isomers and trichloroethylene.

Chromatographic separation for purposes of mass spectral library identification was performed using a 30m X 0.25mm Restek "Stabilwax" capillary column in series with a 100m X 0.25mm J&W DB-1 "Petrocol" capillary column (total length = 130m, both 0.5um film). A 0.1  $\mu l$ volume was injected on-column. The mass spectral scan range was m/z=39-350. The sample was diluted 1:40 with methylene chloride prior to analysis. Many sample peaks identified by mass spectral library search were verified using the following petroleum compound mixtures: n-paraffins (11 compounds), isoparaffins (35 compounds), naphthenes (29 compounds), aromatics (37 compounds) and PNAs (3 compounds). floating product samples were received on May 22, 1997 and stored in the freezer until analysis on June 19-20, 1997.

The methylene chloride core extracts were chromatographed using a 30m X 0.25mm Restek "Stabilwax" (Crossbonded Carbowax-PEG, 0.5µm film) ManTech Environmental Research Services Corporation

capillary column with a SGE Carbowax deactivated Megabore precolumn. SIM mode was used with the ions chosen from those listed in EPA method 524.2 Revision 3.0. Multiple ions were acquired and ion ratios used to verify the accuracy of target compound identification. Standards calibration ranged from 0.025 to 250  $\mu$ g/ml. Complete reports detailing the acquisition method and calibration curves have been recorded. The sample extracts were received on May 22, 1997 and analyzed on June 30, 1997.

If you require further information, please feel free to contact me.

Sincerely

David A. Kovacs

xc: R.L. Cosby

G.B. Smith

J.L. Seeley



Sample	Benzene	TCE	Toluene		X	X-m	징	1,3,5-TMB 1,2,4-TMB 1,2,3-TMB	1,2,4-TMB	1,2,3-TMB
SED-1	Q	3.51E-02	6.22E-02	Q.	2	2.06E-02	2	8	Q	2
SED-2	2	2.41E-02		BLQ	2	BLQ	BLQ	S	2	Q
SED-3	2.08E-02	6.35E-02	9.69E-02	BLQ	2.12E-02	3.58E-02	2.92E-02	2	BLQ	BLQ
SED4	BLQ	6.44E-02	8.77E-02	2	Q	1.79E-02	2	2	2	Q
SED-5	9	7.98E-02	1.26E-01	2	9	3.93E-02	Q	9	S	2
Check Standards (ug/ml)										
20	4.55E+01	0.00E+00	5.99E+01 5.91E+01	5.91E+01	5.95E+01	4.56E+01	5.59E+01	5.95E+01 4.56E+01 5.59E+01 4.60E+01 4.61E+01	4.61E+01	5.22E+01
OC (expected)	5.00E+01	¥X	6.00E+01	6.00E+01	6.00E+01	5.00E+01	6.00E+01	5.00E+01	5.00E+01	5.00E+01
2.5	_	2.41E+00	2.63E+00	2.27E+00	2.63E+00 2.27E+00 2.27E+00 2.29E+00 2.33E+00	2.29E+00	2.33E+00	2.48E+00	2.42E+00	2.55E+00
Method Blank 5-27-97 7.47E-02	7.47E-02	8.03E-01	1.34E+00	BLQ	BLQ	2.31E-01	BLQ		2	<u>Q</u>
MeCl <sub>2</sub> Blank	Q	9	Q	Q	2	Q	Q	2	Q	2

Note: Significant levels of TCE, Toluene and m-Xylene were found in the Method Blank. Sample values may be false positives.



Ref: 97\LB35

June 3, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection and Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198

P.O. Box 1198 Ada, OK 74820

THRU: S.A. Vandegrift 5

Dear Don:

Please find attached the analytical results for Service Request SF-3-266, Shaw AFB, requesting the analysis of ground water samples to be analyzed by purge-and-trap/GC-PID:FID for Benzene, Toluene, Ethylbenzene, p-, m-, & o-Xylene, 1,3,5-, 1,2,4-, & 1,2,3-Trimethylbenzene, and Total Fuel Carbon. We obtained the 55 aqueous samples, in duplicate, in capped, 40 mL VOA autosampler vials May 22, 1997 and they were analyzed May 29-June 2, 1997. The samples were acquired and processed using the Millennium data system. A 6 place (1-1000 ppb) external calibration curve was used to quantitate sample concentration for all the compounds of interest.

RSKSOP-122, "Analysis of Volatile Aromatic Hydrocarbons with Separation of Xylene Isomers by Purge & Trap Gas Chromatography" was used for these analyses. Autosampling was performed using a Dynatech Precision autosampler system in line with a Tekmar LSC 2000 concentrator.

Sincerely,

Lisa R. Black

xc: R.L. Cosby

G.B. Smith

J.T. Wilson

J.L. Seeley

Units = ng/mL

m-XYLENE

P-XYLENE

TOLUENE ETHYLBENZENE

BENZENE

SampleName

		CCCLINE	EINTLBENZENE	p-XYLENE	m-XYLENE	o-XYLENE	1,3,5-TMB	1,2,4-TMB	1,2,3-TMB	Fuel Carbon
IWFFB	95.9	103	103	5	•					
QC, OBSERVED, PPB	48.8	50.6	909	3 :	110	107	116	113	110	V// V
QC, TRUE VALUE, PPB	50.0	000	9.00	<u>×</u> .	53.0	53.5	57.2	56.7	2 4	<b>Y/N</b>
MW1613-1	200	200	0.00	20.0	50.0	50.0	200		J. 1	Z/A
MW1613-1 Duellast	707	BLC	28.0	17.1	B	4.6	9 4	0.00	20.0	A/A
MAN 212 -1 Duplicate	201	BLQ	26.1	, r.	7 6	0.01	15.9	69.5	27.5	924
MVV   613-2	2	2	S	2	ביים מיים	14.4	14.7	62.9	25.9	RR1
MW1613-3	QX	S	2 4	2 :	2	2	2	S	5	3 5
MW1613-3A	Ş	9 9	2 !	2	2	2	S	2	2 4	<b>S</b> :
MW1613-4		2	Q	2	Q	S	2	2 5	2	Q
AAAAA 64 0 4 4	330	203	444	390	243	5 3	€ ;	2	2	ş
A11013-4A	2	2	Q	S	3 2	<b>‡</b> !	เรา	441	175	6470
MW1613-5	1632	984	320	33	2 5	2 !	Q	2	2	2
MW1613-6	Q	Q	Q	\$ 5	i c	427	92.9	295	121	5810
MW1613-/	7.8	26.4	22.4	32.6	5 5	2	Q	Q	2	Q
10 PPB	9.4	10.1	10.5	000	46.2	25.2	33.5	101.7	47.5	2330
MW1613-8	975	570	87.7	2 6	10.5	10.1	10.6	10.4	10.2	<b>4/X</b>
MW1613-9	594	551	Car	40.4	233	151	21.6	75.1	45.9	200
MW1613-9 Duplicate	202	650	2 5	717	331	350	8	22	107	0000
MW1613-9A	S	3 5	212	243	374	412	112	253	140	3830
MW1613-10	2 2	2 5	2	2	2	S	! <u>c</u>	3 4	2 :	4240
MW1613.11	2 5	Q Z	2	Q	S	2	2 9	2 !	2	2
MW/1619-17	1202	1042	38	300	744	9	2 5	2	2	2
MAN/1613-12	58.3	8.5	80.2	60.7	36	9 4	99.0	300	<u>র</u>	6310
MANAGE OF CALINS	BLQ	3.3	2	C	9 C	48.4	45.6	178	99.5	1330
MVV 1613-13	2.5	BLQ	810	2 2	ָרָבָר פּרָבָר	2	2	1.0	BLO	44.3
MW1613-13A	욷	Q	2	ָבָר בְּרַבְּיִר	0.5	BLQ	1.8	1,3	4	7,5
MW1613-14	BLQ	2	2 2	2 9	2	S	2	9	2 2	= 2
500 PPB	465	457	2 5	2	BLQ	2	Q	S	2 2	€ 6
MW1613-14A	S	ş <u>Ş</u>	ş :	451	450	454	455	5 5	Ş §	BLQ.
MW1613-15	Ş	2 2	2	2	BLQ	BLO	S	5 5	ğ :	N/A
MW1613-15A	2 9	2 !	2	2	2	S	2 2	⊋ :	2	BLO
MM/1619 16	2	2	2	Q	2	2 2	ָב ב	S	2	2
MANA 101.0-10	2	S	2	Ş	2 2	2 5	2	2	S	2
MIVI 1013-16 Duplicate	2	Q	9	2	2 6	ב ב	2	2	S	Q
¥ LIN	14.5	943	403	380	ם מ	Ş	Q	2	Q	2
MPB	7.1	917	224	928	27.5	624	졄	321	506	5540
	81.7	1685	242	254	<del>2</del> 6	325	118	326	144	4130
	19.3	159	142	142	9 6	387	118	328	148	4730
ALL INC.	Q	Q	Q	BLO	y C	<u>N</u>	40.4	204	44.9	1720
					,	Ž	Q	2	Q	BLQ

ND = None Detected; BLQ = Below Limit of Quantitation, 1 ppb; N/A = Not Analyzed

3/97 SF-3-266

Fuel Carbon
1,2,3-TMB
1,2,4-TMB
1,3,5-TMB
o-XYLENE
m-XYLENE
p-XYLENE
ETHYLBENZENE
TOLUENE
BENZENE
SampleName

A/N	Y !	2 5	12/	60.4	207	197	529	2	2	2	Q	2	2	Υ/Z	Ϋ́	1.8	16.1	2	930	14.0	2	2	2	2080	2	BLO B	A/Z	623	2350	2	2	2	1290	1280	2	∀X	
45.8	20.0	2	9.7	4.5	BLQ	6.0	27.7	2	Q	9	Q	2	2	2	9.0	2	1.2	2	52.7	Q	오	Q	오	193	2	Q	6.66	20.3	200.	2	2	ջ	41.1	42.7	2	516	
52.0	20.0	2	4.2	2	13.2	13.8	72.3	2	2	2	2	9	2	2	0.8	Q	6.0	Q	137	2.8	2	Q	2	521	2	Q	101	74.9	191	2	2	2	90.5	92.3	2	514	
51.9	50.0	9	6.0	2	BLQ	BLQ	19.0	Q	2	S	2	2	2	Q	0.8	Q	2	Q	3.5	Q	Q	9	2	1771	9	Q	\$	14.3	26.0	2	2	2	13.4	14.0	2	518	) }
49.6	20.0	2	1.0	2.2	2	2	0.6	2	2 5	2 5	2 2	2	2	2	0.8	2	9.1	2	66.7	BLQ	Q	2	2	6	2	2	96.5	59.0	213	2	2	2	123	127	Q	202	}
52,5	20.0	Q	6.0	Q	2	2	40.1	; <u> </u>	2 5	2 5	2 2	2	2	Ş	10	Ę	60	2	65.7	BLO	S	S	2	1037	Q	6.0	106	103	240	2	9	9	190	181	2	787	è
0.70	50.0	Q	2.8	2	8.5	6	25.5	7. C.	2 5	2 5	2 5	2 5	2 5	2 8	0	2 2	-	! <u>S</u>	92.5	2.5	S	9	2	50.5	S	0 18	\$	67.9	135	Q	Q	2	83.5	79.6	S	407	òr
46.8	50.0	2	2.4	S	4	, o	, c	y (2	2 9	2 9	2 9	2 2	2 2	2 2	2 6	? C	2 -	2 5	37.0	97.3	2	2 5	2 5	§ 8	3	<u> </u>	98.7	56.8	113	C	S	S	72.8	74.5	2	2 8	900
47.7	000	Q	80	2	<u> </u>	3 6	7 6	5. C	2 9	2 !	2 9	2 5	2 9	2 2	2 8	0 v	- 6	<u> </u>	<u> </u>	- c	ž 2	2 2	2 5	5 6	3 5	2 5	93.1	122	363	S S	2 5	2 2	208	7 60	i S	2 8	soc .
46.3	2.0	) S	10.5	2 5	1 5	- 10	10.7	28.2	2	Q:	Q !	2 5	2 5	2 5	2 6	ē (	2 5	2.0	2 5	); d	2 4	2 5	2 2	2 8	677	2 2	27.8 87.8	. «	5.5	5 5	2 5	2 2	2 6	90.0	) S	2 !	99
BBB CEVERSON OC	CC, OBSERVED, FFB	dC, INDE VALUE, FFB	1.400	2-1MI	J. UNIT	MT-4	TMP-4 Duplicate	TMP-5	SWS-1	SWS-2	SWS-2 Duplicate	SWS-3	SWS-4	SWS-5	GC LAB BLANK, PPB	1 PPB	TW-120	IW-121	101-701	MW-105	MW-106	70L-WW	MW 107 Duplicate	MW-111	211-WM	MW-112A	MW-113	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	MVV-114	MVV-113	ACT -WM	MW-116	MW-110A	/11-MW	MW 117 Duplicate	MW-118	500 PPB

ND = None Detected; BLQ = Below Limit of Quantitation, 1 ppb; N/A = Not Analyzed



Ref: 97JAD31

June 3, 1997

Dr. Don Kampbell National Risk Management Research Laboratory Subsurface Protection and Remediation Division U.S. Environmental Protection Agency

P.O. Box 1198 Ada, OK 74820

THRU: S.A. Vandegrift

Dear Don:

As requested in Service Request # SF-3-266, headspace GC/MS analysis of 55 Shaw AFB water samples for chlorinated volatile organics was completed. The samples were received on May 22, 1997 and analyzed on May 23-24 & 28-29, 1997. RSKSOP-148 (Determination of Volatile Organic Compounds in Water by Automated Headspace Gas Chromatography/Mass Spectrometry (Saturn II Ion Trap Detector) was used for this analysis.

An internal standard calibration method was established for the 15 compounds. The standard curves were prepared from 1.0 to 4000 ppb. The lower calibration limits were 1.0 ppb.

A dilution corrected quantitation report for the samples, lab duplicates, field duplicates, QC standards and lab blanks is presented in tables 1-3.

If you should have any questions, please feel free to contact me.

Sincerely

John Allen Danie

xc: R.L. Cosby

G.B. Smith

D.D. Fine

J.L. Seeley

J.T. Wilson

Table 1. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

Concentration = ppb

MW1613 8	999   9999999999	MW1613 13A DON ON O
MW1613	99999   99999999	MW 1613 13
MW1613 6	22222222222	1513 ND ND N
MW1613 5	888848888888   8 88884888888888888888888	25 555   5555 5555 5555 5555 5555 5555
MW1613 4A	22222222222   22	MW1613 11 GN
1613 4	22222:222222   2	MW1613 10 10 10 10 10 10 10 10 10 10 10 10 10
1613 3A	2222222222	WW1613 9A OND
1613 3	2222222222	MW1613 90 GN
MW1613 2	2222222222	MW1613 8 ND
MW1613	999999999999	MW1613 Rab Dup ND ND ND ND ND ND ND ND ND ND
Compound	VINYL CHLORIDE  1,1-DICHLOROETHENE  1-1,2-DICHLOROETHENE  1,1-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  CHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE	VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHENE  TRICHLOROETHENE  THICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE

ND = None Detected --- = Below Calibration Limit(1.0 ppb) Dup = Duplicate

Table 2. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

MPC Field Dup	1/10 Dil 31.0 31.0 727 ND 1030 4590 ND ND ND 793 100 11.0 55.0	SWS-2	2222   222222222
MPC	30.4 748 ND 1060 ***** ND ND 718 90.7 90.7 96.2 14.1	SWS-1	2222222222222
MPB	ND 15.4 10.6 11.1 11.3 11.3 11.3 13.5	TMP-5 FT1	1.6 1.8 3.5 42.5 0 NO 0 NO 0 NO 0 NO 0 NO 0 NO 0 NO 0 NO
MPA	0.4.0 0.8.3.9 0.8.5.5.9 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0	TMP-4 FT1	3.5 ND 64.9 169 ND ND ND N
MW1613 16	22222222222222	TMP-3 FT1/0U4	59.6 40.2 ND 1153 150 ND 60.5 ND 2.1 3.9 1.0  ND 6.3 6.3 bove Calibrat
MW1613 15A	22222222222222	TMP-2 OU4 Lab Duo	22.4 113 ND 1030 318 ND 116 ND 1.1 1.1 1.6 5.3 9.8
MW1613 15	2222222222222	TMP-2 OU4	22.8 116 ND 1070 339 ND ND ND 1.2  1.8 5.6 10.5 DII = Dilutio
MW1613 14A		TMP-1 FT-1	ND N
MW1613 14A	999999999999999999999999999999999999999	MPE	AD ND
MW1613 14	999999999999999999999999999999999999999	MPD	ND 1.3 155 1.3 ND 77.0 ND
Compound	VINYL CHLORIDE  1,1-DICHLOROETHENE  1-1,2-DICHLOROETHENE  1,1,1-TRICHLOROETHENE  CARBON TETRACHLORIDE  1,2-DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHENE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  THICHLOROETHENE  THICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,4-DICHLOROBENZENE		VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,5-DICHLOROBENZENE  1,5-DICHLOROBENZE

## Table 3. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

MW 1 0.5.9.00 N N N N N N N N N N N N N N N N N N	MW-115A DN DN D
MW 105 1.2 ND ND N	MW-115 Field Dup 1/10 Dil 474 402 ND 8700 1740  2050 ND ND ND 10.0 ND 10.0 ND 44.0
MF	MW-115 Lab Dup 438 373 ND ***** 1600  2460 ND ND 10.5 4.9 5.1 12.2 47.0
TM-121 FT1 Lab Dup 16.7 7.7 7.7 7.7 80.8 ND ND ND ND ND ND ND ND ND ND ND ND ND	416 368 368 ND 1620  2030 ND ND 10.2 4.8 4.9 11.6 46.4 88.9
TM-121 FT1 15.3 7.3 7.3 80.7 ND ND ND ND ND ND ND ND ND ND ND ND ND	ND 2.1 S8.8 87.1 ND ND N
Field Dup ND ND ND ND ND ND ND ND ND ND ND ND ND	ND N
TT1/0 FT1/0 T10 ON ND ON ON ON ON ON ON ON ON ON ON ON ON ON ON ON ON ON ON	MW-112A ND ND ND ND ND ND ND ND ND ND ND ND ND
SWS-5- QQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQ	ND 2.2 ND 64.8 64.8 22.0 2.3 2.3  ND ND N
SWS P Q Q I I Q Q Q Q Q Q Q Q Q Q Q Q Q Q Q	ND N
SWS-3 D L C C C C C C C C C C C C C C C C C C	MW – 107 ND ND ND 1.1 1.5 ND
Compound  VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  C-1,2-DICHLOROETHENE  C-1,2-DICHLOROETHENE  CHLOROFORM  1,1,1-TRICHLOROETHENE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  THACHLOROETHENE  THACHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE	WWYL CHLORIDE  1,1-DICHLOROETHENE 1,1-DICHLOROETHENE 1,1-DICHLOROETHENE 1,1,1-DICHLOROETHENE 1,1,1-TRICHLOROETHENE 1,2-DICHLOROETHANE 1,2-DICHLOROETHANE 1,2-DICHLOROETHENE 1,3-DICHLOROETHENE 1,4-DICHLOROENZENE 1,4-DICHLOROBENZENE 1,2-DICHLOROBENZENE

## Table 4. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

QC0523D 200 ppb	223 206 210 210 207 212 207 219 222 219	BL0523A BLK	ND N
QC0523C 20 ppb	17.6 22.1 20.3 20.4 20.6 19.5 19.5 20.5 21.4 20.5 21.4 21.8	QC0528M 200 ppb	179 230 208 208 209 202 218 213 206 190 207 208 210 220 bration Limit(4
QC0523B 200 ppb	223 210 205 207 207 207 207 213 213 225 225 225	QC0528L 20 ppb	18.5 23.2 20.4 20.4 20.1 20.5 20.8 20.7 20.9 20.7 19.5 21.0 20.6 = Above Calli
QC0523A 20 ppb	23.3 20.7 20.2 20.2 20.8 20.8 21.1 21.1 22.1 22.1	QC0528K 200 ppb	180 230 201 212 204 201 215 215 203 213 209 219 215 86
MW-118 Lab Dup	999999999999999999999999999999999999999	QC0528J 20 ppb	18.2 23.0 21.2 21.3 21.3 21.8 21.8 22.5 18.3 22.8 20.9 21.5 ol Std. BL =
MW-118	999999  99999999	QC05281 200 ppb	181 230 210 210 216 218 209 209 209 210 217 221 Quality Contro
MW-117 Field Dup	159 159 ND 508 508 508 ND ND 217 25.0 14.0	QC0523H 200 ppb	20.2 25.2 21.8 22.0 20.9 20.4 22.7 22.7 22.3 18.8 23.3 20.2 20.2 20.2 20.1 20.1
MW-117	84.6 148 ND 457 ND ND 203 24.1 2.1 3.4 14.9	QC0523G 20 ppb	18.4 22.6 20.5 21.0 21.1 20.3 20.3 20.3 18.6 19.8 19.6 19.6 19.6
MW-116A	2222222222222222	QC0523F 200 ppb	174 228 210 214 203 203 214 209 210 189 208 216 218 213 221 Jimit(1.0 ppb)
MW-116	9999999999999	QC0523E 20 ppb	18.0 174 21.9 228 20.2 210 21.5 214 20.6 208 20.7 203 20.9 214 20.4 209 22.2 210 17.6 189 19.4 208 21.5 216 18.9 213 20.8 218 20.2 218 20.8 218 20.2 221
Compound	VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CARBON TETRACHLOROETHANE  CARBON TETRACHLORDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE		WINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHENE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  CHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,3-DICHL



Ref: 97-SH33

June 2, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection & Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: S.A. Vandegrift $\zeta \sqrt{}$ 

Dear Don:

Attached are TOC results for 46 Shaw liquids submitted May 28, 1997 under Service Request #SF-3-266. Sample analysis was begun May 29, 1997 and completed June 2, 1997 using RSKSOP-102.

Blanks, duplicates, and AQC samples were analyzed along with your samples, as appropriate, for quality control. If you have any questions concerning this data, please feel free to ask me.

Sincerely,

Sharon Hightower

xc: R.L. Cosby G.B. Smith

J.L. Seeley

### KAMPBELL SHAW LIQUIDS SF-3-266

SAMPLE	MG/L TOC	
TMP-1 TMP-2 TMP-3	1.74 2.85 1.76	•
TMP-4 MW105	2.71 22.2	
MW106 MW107	1.93 1.66	
MW111 MW112 DUP	1.37 22.0 22.0	
MW112A MW113	.884 1.43	
MW114 MW115 MW115A	2.78 8.59 1.06	
MW116 MW117	1.05 3.72	
MW118 MW1613-1 MW1613-2	5.71 1.75 1.52	
MW1613-3 MW1613-3A	.857 .814	
MW1613-4 MW1613-4A MW1613-5	2.59 2.77 50.9	
DUP MW1613-6	51.3 3.82	
MW1613-7 MW1613-8 MW1613-9	1.24 1.88 1.96	
MW1613-9A MW1613-10 MW1613-11	1.27 .740	
MW1613-12 MW1613-13	2.91 4.65 1.30	
MW1613-13A DUP MW1613-14	1.75 1.62 .953	
MW1613-14A MW1613-15	15.7 .932	
MW1613-15A MW1613-16 MPA	2.07 1.24 144	
MPB DUP MPC	5.27 5.77 7.75	
111.0	1.13	

PAGE 2

SAMPLE	MG/L TOC	
MPE 116A TW101	1.51 .909 2.39	
TW121 WSO38	25.5 4.75	

WS038 std t.v.=4.92



May 30, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection & Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: S.A. Vandegrift √

Dear Don:

Attached are TOC results for 5 Shaw soils submitted May 22, 1997 under Service Request #SF-3-266. Sample analysis was begun May 27, 1997 and completed May 30, 1997 using RSKSOP-102 and RSKSOP-120.

Blanks, duplicates, and AQC samples were analyzed along with your samples, as appropriate, for quality control. If you have any questions concerning this data, please feel free to ask me.

Sincerely,

Sharon Hightower

xc: R.L. Cosby

G.B. Smith

J.L. Seeley

KAMPBELL SHAW SOILS SF-3-266

SAMPLE	SOIL FILTRATE % O.C.	SOLIDS % TOC	TOTAL SOIL % TOC	MEAN %TOC	STD DEV
SED 1,1-1 1-2 1-3	.028 .028 .031	.522 .346 .212	.550 .374 .243	.389	.154
SED 2,1-1 1-2 1-3	.058 .066 .070	.956 1.11 1.02	1.01 1.18 1.09	1.09	.085
SED 3,1-1 1-2 1-3	.108 .105 .115	4.88 4.45 4.86	4.99 4.56 4.96	4.84	.240
SED 4,1-1 1-2 1-3	.028 .030 .027	.186 .200 .185	.214 .230 .212	.219	.010
SED 5,1-1 1-2 1-3	.043 .040 .047	10.70 10.42 11.41	10.74 10.46 11.46	10.89	.516
LECO STD WSO38	4.60	.934			

Leco soil std t.v.=.90+/-.04 WSO38 std t.v.=4.92



Ref: 97-BS3/vg 97-MAB6/vg

May 29, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection and Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: Steve Vandegrift

Dear Don:

Please find attached the analytical results for Service Request #SF-3-124 requesting the analysis of 49 field samples for  $\rm CO_2$ , Ferrous Iron, Total Alkalinity, Hydrogen Sulfide, Phenols, and Manganese. Samples were analyzed May 14, 15, 16, 17, 19, and 20, 1997.

Please note that samples with greater than 5.0 mg/l Ferrous Iron may have interfered with the Hach carbon dioxide color test. If you have any questions concerning these results, please feel free to contact us.

Sincerely,

Brad Scroggins

Mark Blankenship

C.B. Cosby G.B. Smith J.L. Seeley Total Alkalinity

	Sample	CO, (mg/L)	Fe <sup>2+</sup> mg/L	(CaCO <sub>3</sub> ) mg/L	H <sub>2</sub> S mg/L	Phenols mg/L
	MW 116μ	28	ND	10	< 0.1	<0.1
	MW 116	18	0.04	20	< 0.1	< 0.1
	TW 101	50	0.84	50	< 0.1	< 0.1
	MW 107	28	1.05	20	< 0.1	< 0.1
	MW 106	32	3.00	20	0.1	< 0.1
	MW 105	88	3.08	20	< 0.1	< 0.1
	MW 111	26	ND	10	< 0.1	< 0.1
	MW 115	84	5.90	70	0.5	< 0.1
	MW 115A	22	0.08	10	< 0.1	< 0.1
	2.0 ppm Std. Fe <sup>2+</sup>		1.94		***	
	MW 118	34	ND	30	< 0.1	< 0.1
	MW 114	36	2.15	20	0.5	< 0.1
	MW 117	44	0.03	. 10	< 0.1	< 0.1
	MW 112	120	7.00	60	0.5	< 0.1
	MW 112A	20	0.01	10	< 0.1	< 0.1
	MW 113	22	0.13	10	< 0.1	< 0.1
	MPC	70	4.00	30	0.3	< 0.1
	MPA	100	4.00	250	0.2	0.94
	MW 1613-6	30	0.12	10	0.1	< 0.1
	MPB	28	3.00	30	2.0	< 0.1
	MW 1613-10	35	0.12	10	< 0.1	< 0.1
	TMP-5	50	4.00	30	< 0.1	< 0.1
	TMP-4	50	4.00	20	0.1	< 0.1
	MW 1613-2	35	0.12	10	<0.1	< 0.1
	MPE	50	7.00	30	< 0.1	< 0.1
	MW 1613-3	30	0.03	20	< 0.1	< 0.1
	TMP-3	40	0.18	10	< 0.1	< 0.1
	TW-121	40	4.00	130	< 0.1	< 0.1
	MW 1613-15	30	0.32	10	< 0.1	< 0.1
	TMP-1	30	0.20	10	< 0.1	< 0.1
	MW 1613-12	115	15.00	50	< 0.1	< 0.1
	TMP-2	80	8.20	20	0.3	< 0.1
	MW 1613-8	80	6.90	90	< 0.1	< 0.1
	MW 1613-9	40	21.00	70	0.1	<0.1
	MW 1613-1	55	14.00	20	< 0.1	< 0.1
	MW 1613-9A	35	ND	10	< 0.1	< 0.1
	MW 1613-11	75	45.00	100	< 0.1	< 0.1
`	-MW 1613-5A	30	0.13	10	< 0.1	< 0.1
	MW 1613-5	70	4.00	100	< 0.1	< 0.1
	MW 1613-7	55	9.00	20	< 0.1	< 0.1
	MW 1613-3A	30	2.01	20	0.5	<0.1

Total Alkalinity

Sample	CO, (mg/L)	$Fe^{2+}$ mg/L	(CaCO <sub>3</sub> ) mg/L	H <sub>2</sub> S mg/L	Phenols mg/L
MW 1613-4	50	8.40	100	<0.1	<0.1
MW 1613-13	70	0.09	10	< 0.1	< 0.1
MW 1613-13A	30	0.05	10	< 0.1	< 0.1
MW 1613-14	55	0.03	10	< 0.1	<0.1
MW 1613-14A	10	0.09	110	0.3	< 0.1
MW 1613-16	35	0.01	30	< 0.1	< 0.1
MW 1613-4A	30	0.03	20	0.3	< 0.1



Ref: 97-MW42/vg 97-LP53/vg

May 29, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection & Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: S.A. Vandegrift

Dear Don:

Attached are the results of 50 Shaw AFB samples submitted to MERSC as part of Service Request #SF-3-266. The samples were received May 22 and analyzed May 23 and 27, 1997. The methods used for analysis were EPA Methods 353.1 for NO<sub>2</sub> and NO<sub>3</sub> and 350.1 for NH<sub>3</sub>, and Waters capillary electrophoresis Method N-601 for Cl and SO<sub>4</sub>. Quality assurance measures performed on this set of samples included spikes, duplicates, known AQC samples and blanks.

If you have any questions concerning these results, please feel free to contact us.

Sincerely,

Mark White

I amda Dannin etan

G.B. Smith
J.L. Seeley

Sample	mg/L Cl	mg/L SO.=	$mg/L NO^{-}_{2} + NO^{-}_{3}(N)$	mg/L NH <sub>3</sub>
MW 1613-1	7.17	<.1	1.89	0.16
MW 1613-2	5.95	<.1	2.40	<.05
MW 1613-3	5.94	<.1	2.80	<.05
MW 1613-3A	5.92	<.1	3.22	<.05
MW 1613-4	5.86	<.1	0.51	1.32
MW 1613-4A	ACID		2.73	<.05
MW 1613-5	5.52	3.56	<.05	1.74
MW 1613-5 Dup	5.39	3.53	<.05	1.75
MW 1613-6	4.76	<.1	3.45	<.05
MW 1613-7	3.58	<.1	0.90	<.05
MW 1613-8	3.89	3.09	<.05	0.91
MW 1613-9	5.70	<.1	0.18	1.16
MW 1613-9 Dup	5.67	<.1		
MW 1613-9A	ACID	IFIED	2.30	<.05
MW 1613-10	4.31	2.36	1.09	<.05
MW 1613-11	4.04	<.1.	<.05	1.33
MW 1613-12	4.68	51.6	0.21	0.47
MW 1613-13	4.07	3.27	0.54	<.05
MW 1613-13A	ACID	IFIED	2.25	<.05
MW 1613-14	2.78	1.27	1.02	<.05
MW 1613-14A	ACID	IFIED	1.84	<.05
MW 1613-15	4.04	17.6	1.51	<.05
MW 1613-15A	ACID	IFIED	1.55	<.05
MW 1613-16	3.05	2.49	1.34	<.05
MW 1613-16 Dup	3.04	2.71		77274
MPA	.58	<.1	<.05	7.41
MPB	<.1	3.21	<.05	1.18
MPC	7.76	<.1	<.05	0.43
MPD	1.07	<.1	<.05	.20
MPE	.35	<.1	<.05	0.17
MPE Dup	.32	<.1	<.05	0.17
TMP-1			IPLE RECEIVED	
TMP-2	ACID		. <.05	0.39
TMP-3	ACID		0.41	<.05
TMP-4	ACID		0.40	0.47
TMP-5		IFIED	<.05	0.37
TW-121	ACID		<.05	6.35
MW-101-TW101	•	ample Receive		0.78
MW-105	3.37	5.57	<.05	2.18
MW-106	.53	<.1	0.11	<.05
MW-107	2.97	4.00	0.26	<.05
MW-111	1.76	<.1	0.59	<.05
MW-112	.52	<.1	<.05	0.83

<u>Sample</u>	mg/L Cl	mg/L SO,=	$mg/L NO^{-}, + NO^{-}(N)$	mg/L NH <sub>3</sub>
MW-112A	2.55	1.53	0.47	<.05
MW-112A Dup	2.53	1.41		
MW-113	3.45	<.1	0.28	<.05
MW-114	5.87	2.75	<.05	<.05
MW-115	20.2	4.30	<.05	1.52
MW-115 Dup			<.05	1.52
MW-115A	3.92	<.1	0.99	<.05
MW-116	3.15	15.1	0.99	<.05
MW-116A	4.26	3.12	0.51	<.05
MW-117	4.23	20.4	<.05	0.26
MW-118	3.08	9.09	0.79	<.05
Unidentified #1	N/A	N/A	0.69	<.05
Unidentified #2	N/A	N/A	0.68	<.05
Unidentified #3	N/A	N/A	1.17	0.76
Blank	<.1	<.1	<.05	<.05
AQC	35.2	44.9	2.05	10.1
*	35.0	44.2		
AQC T.V.	34.8	44.0	2.10	10.0
Spike Rec.	100%	103%	99%	101%
	95%	102%	*****	******

Acidified: An unpreserved sample was not received, prohibiting the analysis of chloride and sulfate.

Sample labels were unreadable on the "unidentified" samples.

Sample	Methane	Ethylene	Ethane
100ppm CH4	91.9	**	***
100ppm C2H4	**	86.6	**
100ppm C2H6	**	**	96.1
HPHe	**	**	-
Lab Blank	**	**	**
TMP-4	0.217	**	**
TMP-5	1.709	*	**
TW-101	0.153	**	**
MW-105	3.594	**	**
MW-106	0.058	***	**
MW-106	0.022	**	**
Lab Dup			
MW-107	0.006	**	**
MW-111	**	**	**
MW-112	6.047	**	**
MW-112A	**	**	**
MW-113	0.001	**	**
MW-113	0.001	**	**
Field Dup			
10ppm CH4	9.0	**	**
MW-114	1.135	**	**
MW-115	2.461	0.005	0.003
MW-115A	**	**	**
MW-116	**	**	**
MW-116A	**	**	**
MW-116A	**	**	**
Lab Dup			
MW-117	0.332	0.003	*
MW-118	0.001	**	**
TW-121	4.541	**	0.010
TW-121	4.164	*	0.012
Field Dup			
100ppm CH4	96.2	**	**
100ppm C2H4	**	86.7	**
100ppm C2H6	**	**	94.4
Lower Limit of C	Quantitation	1	

0.001 0.003 0.002

Units for samples are mg/L.
Units for standards are parts per million.

<sup>\*\*</sup> denotes None Detected.

<sup>\*</sup> denotes Below Limit of Quantitation.

8	-3-200	JIZ0131

	Sample	Methane	Ethylene	Ethane
	100ppm CH4	111.5	**	••
	100ppm C2H4	**	97.8	**
	100ppm C2H6	**	**	91.9
	HPHe	**	**	**
	Lab Blank	**	**	**
	MW1613-1	0.015	**	**
	MW1613-2	**	**	**
	MW1613-3	**	**	**
	MW1613-3A	**	**	**
	MW1613-4	0.072	**	**
	MW1613-4	0.024	**	**
	Lab Dup	3.32.		
	MW1613-4A	•	**	**
	MW1613-5	0.102	**	**
	MW1613-6	**	**	**
	MW1613-7	0.052	**	**
	MW1613-8	0.092	**	**
	MW1613-8	0.092	**	**
	Field Dup	0.032		
	10ppm CH4	9.5	**	**
	MW1613-9	0.009	**	**
	MW1613-9A	•	**	**
	MW1613-10	**	**	**
	MW1613-11	0.138	**	**
	MW1613-12	0.991	-	**
	MW1613-12	0.708	**	**
	Lab Dup			
	MW1613-13	0.019	**	••
	MW1613-13A	0.001	**	**
	MW1613-14	•	**	**
	MW1613-14A	**	**	**
	MW1613-15	**	**	**
	MW1613-15	**	**	**
	Field Dup			
	100ppm CH4	88.2	**	**
	100ppm C2H4	**	99.0	**
	100ppm C2H6	**	**	87.7
	MW1613-15A	**	**	**
	MW1613-16	0.481	**	**
	MPA	6.952	••	**
	MPB	3.378	••	••
	MPC	0.070	•	0.005
	MPC	0.077	•	0.002
	Lab Dup			
	MPD	1.631	••	**
	MPE TMP-1	0.029		-
	•	0.001	-	**
	TMP-2	0.341		
	TMP-3	0.186	Ţ	**
	TMP-3	0.195	•	
	Field Dup 100ppm CH4	04.4	_	-
	00ppm CH4	91.1	00.0	
	00ppm C2H6	**	88.3	90.0
•	Cappill CZMO	W		89.3
L	ower Limit of Qu	antitation		
		0.001	0.003	0.002
				0.002

Units for samples are mg/L.
Units for standards are parts per million.
denotes None Detected.

<sup>\*</sup> denotes Below Limit of Quantitation.



### NATIONAL RISK MANAGEMENT RESEARCH LABORATORY

### **Subsurface Protection and Remediation Division**

(Robert S. Kerr Environmental Research Center)

919 Kerr Research Drive
Post Office Box 1198 Ada, Oklahoma 74820
Fax: 405-436-8703

fax transmittal

to: Todd HerringTon

fAX: 303-831-8208

from:

DON H. KAMPBELL

TEL:405-436-8564

phone:

FAX:405-436-8703

E-Mail:kampbell.donald@epamail.epa.gov

date:

8/11/97

pages:

2 including cover sheet

NOTES: Floating product Analyses

1.2.3-TMB	6.63E+02 1.52E+03 1.27E+03
1.3.5-TMB 1.2.4-TMB	1.74E+03 4.29E+03 3.77E+03
	2.71E+02 4.88E+02 1.7 1.69E+03 1.58E+03 4.28 1.18E+03 1.41E+03 3.7 4,833
m-Xylene o-Xylene	2.71E+02 1.69E+03 1.18E+03
m-Xylene	8.44E+01 3.91E+02 2.8.53E+02 2.28E+03 1.74E+03 1.
p-Xylene	8.44E+01 8.53E+02 7.42E+02
Ethylbenzene	2.45E+02 1.14E+03 8.47E+02
Toluene	ND 3.17E+03 1422.4
TCE	ND 3.35E+03 ND
Benzene	12 BLQ C 2.64E+01 3.3 B ND
Sample	MW1613-12 OU-4 MPC OU-4 MPB
_	

Analyst: D.A. Kovacs

SR #SF-3-266, Shaw AFB, Dr. D. Kampbell

FROM APAB

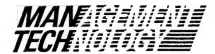
8A98\

Table I. Semi-Quantitative Report for Semi-volatiles found in the Methylene Chloride Extract of Sample MW-112 from Shaw AFB, Service Request SF-3-266.

### Concentration, ng/ml

Benzene	3.4
Toluene	172.8
Ethylbenzene	21.5
m+p-Xylene	814.8
o-Xylene	69.6
1-Methyl-3-ethylbenzene	86.7
1-Methyl-4-ethylbenzene	38.0
1,3,5-Trimethylbenzene	57.9
1-Methyl-2-ethylbenzene	30.3
1,2,4-Trimethylbenzene	178.3
1,2,3-Trimethylbenzene	64.6
Naphthalene	39.9

Pan Pampbell 6/25/97



Ref: 97-MAB5/vg May 27, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection & Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: S.A. Vandegrift  $\leq$ 

Dear Don:

As per Service Request #SFTA-3-125, dissolved hydrogen analysis was performed onsite at Shaw AFB, S.C. using a RGA3 Reduction Gas Analyzer. The analysis began on May 14,1997 and was concluded on May 20, 1997. A six place (0.0 to 10.0PPM) external standard curve was used to quantitate the samples. The following data represents the concentration in the dissolved phase.

	SHAW AFB, S.C.
WELL	H <sub>2</sub> CONCENTRATION (nM)
MW 116	0.38
— MW 116A	6.26
∠ TW 101	1.13
-MW 106	0.14
_ MW 105	0.35
- MW 111	5.79
-MW 115A	0.30
_ MW 115	0.10
MW 118	0.04
MW 117	0.10
- MW 112	3.67
_ MW 112A	19.02

If you have any questions, please see me at your convenience.

Sincerely,

Mark Blankenship

xc: R.L. Cosby G.B. Smith J.L. Seeley



Ref: 97\LB35

June 3, 1997

Dr. Don Kampbell National Risk Management Research Laboratory Subsurface Protection and Remediation Division

U.S. Environmental Protection Agency

P.O. Box 1198

Ada, OK 74820

THRU:

S.A. Vandegrift 5V

Dear Don:

Please find attached the analytical results for Service Request SF-3-266, Shaw AFB, requesting the analysis of ground water samples to be analyzed by purge-and-trap/GC-PID:FID for Benzene, Toluene, Ethylbenzene, p-, m-, & o-Xylene, 1,3,5-, 1,2,4-, & 1,2,3-Trimethylbenzene, and Total Fuel Carbon. We obtained the 55 aqueous samples, in duplicate, in capped, 40 mL VOA autosampler vials May 22, 1997 and they were analyzed May 29-June 2, 1997. The samples were acquired and processed using the Millennium data system. A 6 place (1-1000 ppb) external calibration curve was used to quantitate sample concentration for all the compounds of interest.

RSKSOP-122, "Analysis of Volatile Aromatic Hydrocarbons with Separation of Xylene Isomers by Purge & Trap Gas Chromatography" was used for these analyses. Autosampling was performed using a Dynatech Precision autosampler system in line with a Tekmar LSC 2000 concentrator.

Sincerely,

Lisa R. Black

xc: R.L. Cosby

G.B. Smith

J.T. Wilson

J.L. Seeley

David Kovacs

Black

SamolaNama	RENZENE	TOLUENE	ETHYLBENZENE	p-XYLENE	m-XYLENE	o-XYLENE	1,3,5-TMB	1,2,4-TMB	1,2,3-TMB	Fuel Carbon
	c c	5	9	100	110	107	116	113	110	ΑX
100 PPB	B.C.B.	3 6	200	24.7	53.0	53.5	57.2	26.7	49.3	∢ Z
OC, OBSERVED, PPB	48.8 50.0	20.0	9,00	; ç	2.05	50.0	50.0	20.0	20.0	Ϋ́Z
QC, TRUE VALUE, PPB	50.0	0.00	0.00	42.5	2 6	15.5	15.9	69.5	27.5	924
MW1613-1	707	ָ ק ק	26.0	- u	7 0	14.4	14.7	62.9	25.9	881
MW1613-1 Duplicate	5 5	ָרָ בְּיֵלְ בְּיִלְ	1,07	2	Š	C	2	2	2	2
MW1613-2	2 9	2 9	2 5	2 5	2	2	2	Q	2	2
MW1613-3	2 9	2 5	2 2	2 2	Ş	S	2	2	2	2
MW1613-3A	2 5	2 8	5 5	<u> </u>	543	44	151	441	175	6470
MW1613-4	35. 13.	3 5	£ 2	3 5	3 2	2	9	2	Q	Q
MW1613-4A	5 5	2 8	330	8 8	551	427	92.9	295	121	5810
MW1613-5	7 0	\$ 2	3 5	Ş	S	2	9	9	Q	Q
MW1613-6	Z 8	£ %	22.4	23.5	48.2	25.2	33.5	101.7	47.5	2330
MW1613-7	9. 6	101	105	10.3	10.5	10.1	10.6	10.4	10.2	A/N
10 FFB	4.0	2.2	7.78	96.4	233	151	21.6	75.1	45.9	3060
MW1613-8	6/6	2 4	180	217	331	350	100	222	127	3930
MW1613-9	<b>T</b> 8	100	3 6	543	374	412	112	253	143	4240
MW1613-9 Duplicate	¥ 5	200	2 4	2 5	<u> </u>	Ş	2	2	2	2
MW1613-9A	2 5	2 2	2 9	2 2	2 5	2	S	2	2	2
MW1613-10	2	2	2 8	2 8	2 %	486	000	300	183	6310
MW1613-11	1202	1042	500	20 1	¥ ;	8 6	20.00	478	90 8	1330
MW1613-12	58.3	8.5	80.2	60.7	380	/8.4 	9.0	2 -	. C	44.3
MW1613-12rinse	BLQ	3.3	2	2	BLQ	2	2 :	0. 9	הרב י	7.4.4
MW1613-13	2.5	BLQ	BLO	BLQ	1.0	BLQ	9.	E		= 4
MW1613-13A	2	2	2	9	2	2	2	2 9	2 :	2 2
MW1613-14	BLQ	9	2	2	BLQ	2	2	2	2	
500 PPB	465	457	452	451	450	454	455	<u>4</u>	94 9	<b>∀</b> (2)
MW1613-14A	9	2	2	2	BLQ	BLQ	2	2	2	o e
MW1613-15	2	2	2	9	2	2	9	2	2	2 :
MW1613-15A	2	9	2	g	2	2	2	9	2	2
MM/1613-16	Ş	2	2	2	2	2	ᄝ	2	2	2
MIN 1013-10 MAN/1613-16 Direlicate	S	2	2	2	BLQ	9	2	2	2	ВГО
MPA CIS-10 Cupicato	14.5	943	403	388	922	624	52	321	506	5540
MON	7.1	917	224	228	548	325	118	326	4	4130
O LA	81.7	1685	242	254	638	387	118	328	148	4730
MPO	19.3	159	142	142	232	122	40.4	204	44.9	1720
	2	S	CZ	C	C	2	Q	2	2	8 0
T L	2	2	2			ļ				

SampleName	BENZENE	TOLUENE	ETHYLBENZENE	p-XYLENE	m-XYLENE	o-XYLENE	1,3,5-TMB	1,2,4-TMB	1,2,3-TMB	Fuel Carbon
OC OBSERVED PPR	46.3	47.7	46.8	54.0	52.5	49.6	51.9	52.0	45.8	Y/X
OC TRUE VALUE PPR	50.0	50.0	20.0	50.0	50.0	20.0	50.0	50.0	20.0	₹ Z
TMP.1	S	2	2	Q	2	2	2	2	2	2
TMP-2	12.5	2.3	2.4	2.8	6.0	1.0	6.0	4.2	9.7	127
TMP-3	12.4	9	2	Q	2	2.2	2	2	4.5	60.4
TMP-4	10.1	BLO	8.4	8.5	9	2	BLQ	13.2	BLQ	207
TMP-4 Dinifcate	10.7	BLO	8.9	8.3	Q	Q	BLQ	13.8	6.0	197
TMP.5	28.2	6.6	77.2	70.2	49.1	9.0	19.0	72.3	27.7	529
SWS-1	2	2	Q	2	Q	Q	Q	2	9	2
c-sws	9	2	9	Q	Q	2	2	2	2	2
SWS-2 Duolicate	2	2	2	9	2	Q	Q	9	Q	2
SWS-3	2	2	2	2	Q	Q	Q	2	2	2
5WS-4	2	2	2	Q	9	2	Q	2	9	2
SWS-5	2	2	2	2	Q	Q	Q	잎	Q	2
GC LAB BLANK, PPB	2	2	2	2	9	2	g	9	9	Y.
1 PPB	6.0	0.8	6.0	6.0	1.0	0.8	0.8	0.8	0.8	Ϋ́Z
TW-120	9	2.1	2	2	9	9	2	9	2	4.8
TW-121	5.3	1.2	1.0	1.2	0.9	1.9	Q	6.0	1.2	16.1
101-WT	2	2	Q	Q	Q	Q	Q	9	2	2
MW-105	70.7	7.1	37.9	92.5	65.7	66.7	3.5	137	52.7	930
MW-106	BLO	BLQ	2.6	2.1	BLQ	BLQ	Q	2.8	2	14.0
MW-107	2	2	Q	2	9	9	9	운	2	2
MW 107 Duplicate	2	2	Q	2	9	Q	9	2	2	2
MW-111	Q	Q	2	2	9	2	9	2	2	2
MW-112	229	438	495	501	1037	199	171	<u>8</u> 2	193	2080
MW-112A	2	2	2	Q	Q	9	9	2	2	2
MW-113	Q	Q	BLQ	BLQ	6.0	9	2	2	2	BLQ
100 PPB	87.8	93.1	98.7	104	105	96.5	\$	101	99.9	¥ S
MW-114	6.3	12.2	56.8	57.9	<del>1</del>	29.0	14.3	74.9	20.3	623
MW-115	157	393	113	135	240	213	26.0	191	20.7	7320
MW-115A	2	Q	Q	Q	오	2	2	2	2	2 !
MW-116	9	Q	2	9	9	2	2	2	2	2 !
MW-116A	Q	Q	Q	2	Q	2	2	2	2	2
MW-117	30.0	89.5	72.8	83.5	<del>1</del>	123	13.4	90.5	41.1	0621
MW 117 Duplicate	30.7	93.7	74.5	9.62	181	127	14.0	92.3	42.7	1280
MW-118	9	Q	2	9	2	2	2	2	2	2
500 PPB	499	503	206	487	487	200	518	514	516	ď Z

ND = None Detected; BLQ = Below Limit of Quantitation, 1 ppb; N/A = Not Analyzed



Ref: 97JAD31

June 3, 1997

Dr. Don Kampbell National Risk Management Research Laboratory Subsurface Protection and Remediation Division U.S. Environmental Protection Agency P.O. Box 1198 Ada, OK 74820

THRU: S.A. Vandegrift N

### Dear Don:

As requested in Service Request # SF-3-266, headspace GC/MS analysis of 55 Shaw AFB water samples for chlorinated volatile organics was completed. The samples were received on May 22, 1997 and analyzed on May 23-24 & 28-29, 1997. RSKSOP-148 (Determination of Volatile Organic Compounds in Water by Automated Headspace Gas Chromatography/Mass Spectrometry (Saturn II Ion Trap Detector) was used for this analysis.

An internal standard calibration method was established for the 15 compounds. The standard curves were prepared from 1.0 to 4000 ppb. The lower calibration limits were 1.0 ppb.

A dilution corrected quantitation report for the samples, lab duplicates, field duplicates, QC standards and lab blanks is presented in tables 1-3.

If you should have any questions, please feel free to contact me.

Sincerely

John Allen Danie

xc: R.L. Cosby

G.B. Smith

D.D. Fine

J.L. Seeley

J.T. Wilson

## Table 1. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

Concentration = ppb

Compound	MW1613	MW1613	1613 3	1613 3A	1613 4	MW1613 4A	MW1613 5	MW1613 6	MW1613	MW1613 8
VINYL CHLORIDE  1,1-DICHLOROETHENE  1-1,2-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROBENZENE  1,3-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE	999999999999	2222222222111		222222222111	99999:9999999	999999999999999999999999999999999999999	55555;5555555	999999999999   9	99999 9999999	222   222222222
VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  CHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE	MW1613  Rab Dup  ND  ND  ND  ND  ND  ND  ND  ND  ND  N	Field Dup ND ND ND ND ND ND ND ND ND ND ND ND ND	MW 1613	WW1613 9A ON	MW1613 10 10 10 10 10 10 10 10 10 10 10 10 10	MW1613 1- GN	25	1613 ND ND N	MW1613 13 13 13 13 13 13 13 13 13 13 13 13 1	MW1613 134 A L L L L L L L L L L L L L L L L L L

ND = None Detected --- = Below Calibration Limit(1.0 ppb) Dup = Duplicate

# Table 2. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

MPC Field Dup	31.0 31.0 1030 4590 ND ND 23900 ND ND 793 100 11.0 55.0	SWS-2	2222   222222222
MPC	30.4 748 ND 1060 ***** ND ND 718 90.7 90.7 96.2 14.1 68.2	SWS1	2222222222222
MPB	ON 15.4 10.6 10.6 10.6 10.6 10.6 10.6 10.6 10.6	TMP-5 FT1	116 00 00 00 00 00 00 00 00 00 00 00 00 00
MPA	ON 4.3 3.5 4.0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	TMP-4 FT1	7.7 3.5 ND 64.9 169 ND ND ND ND ND 2.2 2.2 2.5 tion Limit(40
MW1613 16	999999999999999999999999999999999999999	TMP-3 FT1/0U4	59.6 40.2 ND 1153 150 ND 60.5 ND 2.1 3.9 1.0  ND 6.3 6.3 6.3
MW1613 15A	2222222222222	TMP-2 004	22.4 113 1030 318 318 ND 116 1.1 1.6 5.3 9.8 9.8
MW1613 15	2222222222222	TMP-2 OU4	22.8 116 ND 339 ND 78.6 ND ND 1.2  1.8 5.6 10.5 Dilutic
MW1613		TMP-1 FT-1	Limit(1.0 ppb)
MW1613 14A	99999999999999	MPE	17 ND ND 1.4 1.4 1.4 1.4 1.4 1.4 1.4 1.4 1.4 1.4
MW1613 14	2222222222222	MPD	ND
Compound	VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  CHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE		WINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROBENZENE  1,3-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE

## Table 3. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

MW106	5   5 % % S S S S S S S S S S S S S S S S S	MW-115A	525   55555   55555
MW 105	2.1 - 0	MW-115 Field Dup	474 402 ND 8700 1740 1740 ND ND 10.0 ND 11.0 44.0
TMP-101	99999999999999	MW-115 Lab Dup	438 373 ND 1600  ND ND 10.5 4.9 5.1 12.2 47.0
TM-121 FT1	Lab Dup 16.7 7.7 2.2 30.8 30.8 ND ND ND ND ND ND 1.6	MW-115	416 368 ND 1620  2030 ND ND 10.2 4.8 4.9 4.9 4.9 4.9 46.4 88.9
TM-121 FT1	15.3 7.3 1.5 1.5 30.7 ND ND ND ND ND ND ND ND ND ND ND ND ND	MW-114	ND S8.8 87.1 ND ND ND ND ND ND ND ND ND ND ND ND ND
TM-120 FT1/0U4		MW-113	ND 1.2 1.2 ND ND ND ND ND ND ND ND ND ND ND ND ND
TM-120 FT1/0U4		MW-112A	ON O
SWS5	999999999999999999999999999999999999999	MW-112	ND 2.2 ND 64.8 22.0 22.0 22.3 ND
SWS-4	999     9999999999	MW-111	ND N
SWS-3	S   S = 25   S S S S S S S S S S S S S S S S S S	MW-107	ND ND HIS TO A STATE OF THE STA
Compound	WNYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2-DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHANE  1,2-DICHLOROETHENE  CHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE		WINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  C-1,2-DICHLOROETHENE  CARBON TETRACHLORIDE  1,2-DICHLOROETHANE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  THICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,5-DICHLOROBENZENE  1,5-DICHLOROBE

# Table 4. Quantitation Report for S.R. # SF-3-266 from Shaw AFB.

Compound	MW-116	MW-116A	MW-117	MW-117 Field Dup	MW-118	MW-118 Lab Dup	QC0523A 20 ppb	QC0523B 200 ppb	QC0523C 20 ppb	QC0523D 200 ppb
VINYL CHLORIDE  1,1 – DICHLOROETHENE  T – 1,2 – DICHLOROETHENE  1,1 – DICHLOROETHANE  CHLOROFORM  1,1,1 – TRICHLOROETHANE  CARBON TETRACHLORIDE  1,2 – DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHENE  TRICHLOROETHENE  TRICHLOROETHENE  THACHLOROETHENE  THACHLOROBENZENE  1,3 – DICHLOROBENZENE  1,4 – DICHLOROBENZENE  1,2 – DICHLOROBENZENE  1,2 – DICHLOROBENZENE	22222222222222	2222222222222	84.6 148 ND 481 457  ND ND 203 24.1 2.1 3.4 14.9	94.0 159 ND 509 508 ND 4690 ND ND 217 25.0 14.0	22222   2222222	999999   99999999	23.3 20.7 20.2 20.2 20.8 20.8 21.1 21.1 22.1 22.1 22.1	223 205 207 207 207 207 208 213 225 225 225	17.6 22.1 20.3 20.4 20.4 20.6 19.5 19.5 20.4 20.5 21.7 21.7	223 206 210 210 214 207 212 204 222 218 222 219 219
	QC0523E 20 ppb	QC0523F 200 ppb	QC0523G 20 ppb	QC0523Н 200 ррb	QC05281 200 ppb	QC0528J 20 ppb	QC0528K 200 ppb	QC0528L 20 ppb	QC0528M 200 ppb	BL0523A BLK
VINYL CHLORIDE  1,1-DICHLOROETHENE  1,1-DICHLOROETHENE  1,1-DICHLOROETHANE  C-1,2-DICHLOROETHANE  CHLOROFORM  1,1,1-TRICHLOROETHANE  1,2-DICHLOROETHANE  TRICHLOROETHANE  TRICHLOROETHENE  TETRACHLOROETHENE  TETRACHLOROETHENE  THICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,2-DICHLOROBENZENE  1,3-DICHLOROBENZENE  1,4-DICHLOROBENZENE  1,5-DICHLOROBENZENE  1,5-DICHLOROBENZEN	18.0 174 21.9 228 20.2 210 21.5 214 20.6 208 20.7 203 20.9 214 20.4 209 22.2 210 17.6 189 19.4 208 21.5 216 18.9 213 20.8 213 20.8 218 20.2 221	174 228 210 214 203 203 214 209 210 189 208 216 218 221 Limit(1.0 ppb)	18.4 22.6 20.5 21.0 21.1 20.3 20.3 20.3 18.0 19.8 19.8 19.6 19.6	20.2 25.2 21.8 22.0 20.9 20.4 22.7 22.3 18.8 22.3 22.3 20.2 20.2 20.6 20.1 ficate QC =	181 230 210 216 216 218 209 209 209 210 217 221 221 Quality Cont	18.2 23.0 21.2 21.5 21.5 21.3 22.5 18.3 22.8 22.8 20.6 20.9 21.5	180 230 212 212 204 204 215 215 209 213 219 219 219	18.5 23.2 20.4 20.1 20.5 20.8 20.7 20.8 17.8 20.9 20.7 19.5 20.9 20.7	179 230 208 209 199 202 218 207 207 208 210 220 Ilibration Limit	ND N



Ref: 97-SH33

June 2, 1997

Dr. Don Kampbell National Risk Management Research Laboratory Subsurface Protection & Remediation Division U.S. Environmental Protection Agency P.O. Box 1198 Ada, OK 74820

THRU: S.A. Vandegrift (

Dear Don:

Attached are TOC results for 46 Shaw liquids submitted May 28, 1997 under Service Request #SF-3-266. Sample analysis was begun May 29, 1997 and completed June 2, 1997 using RSKSOP-102.

Blanks, duplicates, and AQC samples were analyzed along with your samples, as appropriate, for quality control. If you have any questions concerning this data, please feel free to ask me.

Sincerely,

Sharon Hightower

xc: R.L. Cosby G.B. Smith

J.L. Seeley

KAMPBELL SHAW LIQUIDS SF-3-266

SAMPLE	MG/L	TOC
TMP-1 TMP-2 TMP-3 TMP-4 MW105 MW106 MW107 MW111 MW112 DUP MW112A MW113 MW114 MW115 MW115A MW116 MW117 MW118 MW1613-1 MW1613-2 MW1613-3 MW1613-3 MW1613-4 MW1613-4 MW1613-4 MW1613-5 DUP MW1613-6 MW1613-7 MW1613-7 MW1613-8	1.74 2.85 1.76 2.71 22.93 1.66 1.37 22.0 1.66 1.37 22.0 1.75 1.75 1.75 1.75 2.77 1.75 2.77 1.75 2.77 1.85 7.79 1.88	TOC
MW1613-8 MW1613-9 MW1613-10 MW1613-11 MW1613-12 MW1613-13 MW1613-13A DUP MW1613-14 MW1613-14A MW1613-15 MW1613-15A MW1613-16 MPA MPB DUP MPC	1.88 1.96 1.27 .740 2.91 4.65 1.75 1.62 .953 15.7 .932 2.07 1.24 15.27 7.75	

PAGE 2

SAMPLE	MG/L TOC	
MPE 116A TW101 TW121 WSO38	1.51 .909 2.39 25.5 4.75	

WS038 std t.v.=4.92



May 30, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection & Remediation Division
U.S. Environmental Protection Agency

P.O. Box 1198 Ada, OK 74820

THRU: S.A. Vandegrift √

Dear Don:

Attached are TOC results for 5 Shaw soils submitted May 22, 1997 under Service Request #SF-3-266. Sample analysis was begun May 27, 1997 and completed May 30, 1997 using RSKSOP-102 and RSKSOP-120.

Blanks, duplicates, and AQC samples were analyzed along with your samples, as appropriate, for quality control. If you have any questions concerning this data, please feel free to ask me.

Sincerely,

Sharon Hightower

xc: R.L. Cosby

G.B. Smith

J.L. Seeley

KAMPBELL SHAW SOILS SF-3-266

SAMPLE	SOIL FILTRATE % O.C.	SOLIDS % TOC	TOTAL SOIL % TOC	MEAN %TOC	STD DEV
SED 1,1-1 1-2 1-3	.028 .028 .031	.522 .346 .212	.550 .374 .243	.389	.154
SED 2,1-1 1-2 1-3	.058 .066 .070	.956 1.11 1.02	1.01 1.18 1.09	1.09	.085
SED 3,1-1 1-2 1-3	.108 .105 .115	4.88 4.45 4.86	4.99 4.56 4.96	4.84	.240
SED 4,1-1 1-2 1-3	.028 .030 .027	.186 .200 .185	.214 .230 .212	.219	.010
SED 5,1-1 1-2 1-3	.043 .040 .047	10.70 10.42 11.41	10.74 10.46 11.46	10.89	.516
LECO STD WSO38	4.60	.934			

Leco soil std t.v.=.90+/-.04 WSO38 std t.v.=4.92



Ref: 97-BS3/vg 97-MAB6/vg

May 29, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection and Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: Steve Vandegrift 5

Dear Don:

Please find attached the analytical results for Service Request #SF-3-124 requesting the analysis of 49 field samples for CO<sub>2</sub>, Ferrous Iron, Total Alkalinity, Hydrogen Sulfide, Phenols, and Manganese. Samples were analyzed May 14, 15, 16, 17, 19, and 20, 1997.

Please note that samples with greater than 5.0 mg/l Ferrous Iron may have interfered with the Hach carbon dioxide color test. If you have any questions concerning these results, please feel free to contact us.

Sincerely,

**Brad Scroggins** 

Mark Blankenship

xc: R.L. Cosby G.B. Smith J.L. Seeley Total Alkalinity

Sample	CO, (mg/L)	Fe <sup>2+</sup> mg/L	(CaCO <sub>3</sub> ) mg/L	H <sub>2</sub> S mg/L	Phenols mg/L
MW 116μ	28	ND	10	< 0.1	< 0.1
MW 116	18	0.04	20	< 0.1	< 0.1
TW 101	50	0.84	50	< 0.1	< 0.1
MW 107	28	1.05	20	< 0.1	< 0.1
MW 106	32	3.00	20	0.1	< 0.1
MW 105	88	3.08	20	< 0.1	< 0.1
MW 111	26	ND	10	< 0.1	< 0.1
MW 115	84	5.90	70	0.5	< 0.1
MW 115A	22	0.08	10	< 0.1	< 0.1
2.0 ppm Std. Fe <sup>2+</sup>		1.94			
MW 118	34	ND	30	< 0.1	< 0.1
MW 114	36	2.15	20	0.5	< 0.1
MW 117	44	0.03	10	< 0.1	< 0.1
MW 112	120	7.00	60	0.5	< 0.1
MW 112A	20	0.01	10	< 0.1	< 0.1
MW 113	22	0.13	10	< 0.1	< 0.1
MPC	70	4.00	30	0.3	< 0.1
MPA	100	4.00	250	0.2	0.94
MW 1613-6	30	0.12	10	0.1	< 0.1
MPB	28	3.00	30	2.0	< 0.1
MW 1613-10	35	0.12	10	< 0.1	< 0.1
TMP-5	50	4.00	30	< 0.1	< 0.1
TMP-4	50	4.00	20	0.1	< 0.1
MW 1613-2	35	0.12	. 10	< 0.1	< 0.1
MPE	50	7.00	30	< 0.1	<0.1
MW 1613-3	30	0.03	20	< 0.1	<0.1
TMP-3	40	0.18	10	<0.1	<0.1
TW-121	40	4.00	130	<0.1	<0.1
MW 1613-15	30	0.32	10	<0.1	<0.1
TMP-1	30	0.20	10	<0.1	<0.1
MW 1613-12	115	15.00	50	<0.1	<0.1
TMP-2	80	8.20	20	0.3	<0.1
MW 1613-8	80	6.90	90	<0.1	<0.1
MW 1613-9	40	21.00	70	0.1	<0.1
MW 1613-1	55	14.00	20	<0.1	<0.1
MW 1613-9A	35	ND	10	<0.1	<0.1
MW 1613-11	75	45.00	100	<0.1	<0.1
MW 1613-5A	30	0.13	10	<0.1	<0.1
MW 1613-5	70 55	4.00	100	<0.1	<0.1
MW 1613-7	55	9.00	20	<0.1	<0.1
MW 1613-3A	30	2.01	20	0.5	< 0.1

Total Alkalinity

Sample	<u>CO</u> , (mg/L)	Fe <sup>2+</sup> mg/L	(CaCO <sub>3</sub> ) mg/L	H <sub>2</sub> S mg/L	Phenols mg/L
MW 1613-4	50	8.40	100	< 0.1	<0.1
MW 1613-13	70	0.09	10	< 0.1	< 0.1
MW 1613-13A	30	0.05	10	< 0.1	< 0.1
MW 1613-14	55	0.03	10	< 0.1	< 0.1
MW 1613-14A	10	0.09	110	0.3	< 0.1
MW 1613-16	35	0.01	30	< 0.1	< 0.1
MW 1613-4A	30	0.03	20	0.3	< 0.1



Ref: 97-MW42/vg 97-LP53/vg

May 29, 1997

Dr. Don Kampbell
National Risk Management Research Laboratory
Subsurface Protection & Remediation Division
U.S. Environmental Protection Agency
P.O. Box 1198
Ada, OK 74820

THRU: S.A. Vandegrift

Dear Don:

Attached are the results of 50 Shaw AFB samples submitted to MERSC as part of Service Request #SF-3-266. The samples were received May 22 and analyzed May 23 and 27, 1997. The methods used for analysis were EPA Methods 353.1 for NO<sub>2</sub> and NO<sub>3</sub> and 350.1 for NH<sub>3</sub>, and Waters capillary electrophoresis Method N-601 for Cl and SO<sub>4</sub>. Quality assurance measures performed on this set of samples included spikes, duplicates, known AQC samples and blanks.

If you have any questions concerning these results, please feel free to contact us.

Sincerely,

Mark White

Lynda Pennington

G.B. Smith
J.L. Seeley

Sample	mg/L Cl	mg/L SO.	$mg/L NO_2 + NO_3(N)$	mg/L NH
MW 1613-1	7.17	<.1	1.89	0.16
MW 1613-2	5.95	<.1	2.40	<.05
MW 1613-3	5.94	<.1	2.80	<.05
MW 1613-3A	5.92	<.1	3.22	<.05
MW 1613-4	5.86	<.1	0.51	1.32
MW 1613-4A	ACID		2.73	<.05
MW 1613-5	5.52	3.56	<.05	1.74
MW 1613-5 Dup	5.39	3.53	<.05	1.75
MW 1613-6	4.76	<.1	3.45	<.05
MW 1613-7	3.58	<.1	0.90	<.05
MW 1613-8	3.89	3.09	<.05	0.91
MW 1613-9	5.70	<.1	0.18	1.16
MW 1613-9 Dup	5.67	<.1	0.16	1.10
MW 1613-9 Dup		IFIED	2.30	<.05
MW 1613-10	4.31	2.36	1.09	<.05
MW 1613-11	4.04	<.1.	<.05	1.33
MW 1613-11				
	4.68	51.6	0.21	0.47
MW 1613-13	4.07	3.27	0.54	<.05
MW 1613-13A		IFIED	2.25	<.05
MW 1613-14	2.78	1.27	1.02	<.05
MW 1613-14A		IFIED	1.84	<.05
MW 1613-15	4.04	17.6	1.51	<.05
MW 1613-15A		IFIED	1.55	<.05
MW 1613-16	3.05	2.49	1.34	<.05
MW 1613-16 Dup	3.04	2.71	100 mm one one	
MPA	.58	<.1	<.05	7.41
MPB	<.1	3.21	<.05	1.18
MPC	7.76	<.1	<.05	0.43
MPD	1.07	<.1	<.05	.20
MPE	.35	<.1	<.05	0.17
MPE Dup	.32	<.1	<.05	0.17
TMP-1			IPLE RECEIVED	
TMP-2		IFIED	<.05	0.39
TMP-3		IFIED	0.41	<.05
TMP-4		IFIED	0.40	0.47
TMP-5		IFIED	<.05	0.37
TW-121		IFIED	<.05	6.35
MW-101		Sample Receive		0.78
MW-105	3.37	5.57	<.05	2.18
MW-106	.53	<.1	0.11	<.05
MW-107	2.97	4.00	0.26	<.05
MW-111	1.76	<.1	0.59	<.05
MW-112	.52	<.1	<.05	0.83

Sample	mg/L Cl	mg/L SO,=	$mg/L NO^{\circ}_{2} + NO^{\circ}_{3}(N)$	mg/L NH <sub>3</sub>
MW-112A	2.55	1.53	0.47	<.05
MW-112A Dup	2.53	1.41		
MW-113	3.45	<.1	0.28	<.05
MW-114	5.87	2.75	<.05	<.05
MW-115	20.2	4.30	<.05	1.52
MW-115 Dup			<.05	1.52
MW-115A	3.92	<.1	0.99	<.05
MW-116	3.15	15.1	0.99	<.05
MW-116A	4.26	3.12	0.51	<.05
MW-117	4.23	20.4	<.05	0.26
MW-118	3.08	9.09	0.79	<.05
Unidentified #1	N/A	N/A	0.69	<.05
Unidentified #2	N/A	N/A	0.68	<.05
Unidentified #3	N/A	N/A	1.17	0.76
Blank	<.1	<.1	<.05	<.05
AQC	35.2	44.9	2.05	10.1
	35.0	44.2	****	
AQC T.V.	34.8	44.0	2.10	10.0
Spike Rec.	100%	103%	99%	101%
	95%	102%		

Acidified: An unpreserved sample was not received, prohibiting the analysis of chloride and sulfate.

Sample labels were unreadable on the "unidentified" samples.

Sample	Methane	Ethylene	Ethane
100ppm CH4	91.9	***	***
100ppm C2H4	**	86.6	**
100ppm C2H6	Web.	**	96.1
HPHe	**	**	**
Lab Blank	**	**	**
TMP-4	0.217	**	***
TMP-5	1.709	•	**
TW-101	0.153	**	***
MW-105	3.594	**	**
MW-106	0.058	**	***
MW-106	0.022	**	**
Lab Dup			
MW-107	0.006	**	**
-MW-111	**	**	de de
MW-112	6.047	**	**
MW-112A	**	**	**
MW-113	0.001	**	**
MW-113	0.001	**	**
Field Dup			
10ppm CH4	9.0	**	***
MW-114	1.135	**	**
MW-115	2.461	0.005	0.003
MW-115A	**	trier	***
MW-116	**	100	and the
MW-116A	**	**	**
MW-116A	**	No.	**
Lab Dup			
MW-117	0.332	0.003	*
MW-118	0.001	**	**
TW-121	4.541	***	0.010
TW-121	4.164	•	0.012
Field Dup			
100ppm CH4	96.2	**	**
100ppm C2H4	***	86.7	
100ppm C2H6	***	**	94.4
Lower Limit of 0	Quantitation	1	

0.001 0.003 0.002

Units for samples are mg/L.
Units for standards are parts per million.

<sup>\*\*</sup> denotes None Detected.

\* denotes Below Limit of Quantitation.

Sample	Methane	Ethylene	Ethane
100ppm CH4	111.5	**	**
100ppm C2H4	**	97.8	**
100ppm C2H6	**	**	91.9
HPHe	**	**	**
Lab Blank	**	**	**
MW1613-1	0.015	**	**
MW1613-2	••	**	**
MW1613-3	**	**	**
MW1613-3A	***	**	**
MW1613-4	0.072	**	**
MW1613-4	0.072	**	**
	0.024		
Lab Dup		**	
MW1613-4A			
MW1613-5	0.102	**	**
MW1613-6	**		
MW1613-7	0.052	***	**
MW1613-8	0.092	**	**
MW1613-8	0.092	**	**
Field Dup			
10ppm CH4	9.5		**
MW1613-9	0.009	**	**
MW1613-9A	•	**	**
MW1613-10	**	**	**
MW1613-11	0.138	**	**
MW1613-12	0.991	**	**
MW1613-12	0.708	**	**
Lab Dup	0 00		
MW1613-13	0.019	**	**
MW1613-13A	0.001	**	**
MW1613-14	0.001	**	**
MW1613-14A	**	**	**
MW1613-15		••	**
MW1613-15	**	**	**
Field Dup			
100ppm CH4	88.2	••	••
100ppm C2H4	00.2	00.0	**
100ppm C2H4	••	99.0	87.7
		**	87.7
MW1613-15A MW1613-16	0.404	**	**
MPA	0.481	**	**
MPB	6.952	••	
	3.378	•	2 225
MPC	0.070	-	0.005
MPC	0.077	•	0.002
Lab Dup			
MPD	1.631		
MPE	0.029	**	**
TMP-1	0.001	-	
TMP-2	0.341	•	44
TMP-3	0.186	•	**
TMP-3	0.195	•	**
Field Dup			
100ppm CH4	91.1	**	**
100ppm C2H4	••	88.3	**
100ppm C2H6	**	**	89.3
ower Limit of C			
	0.001	0.003	0.002

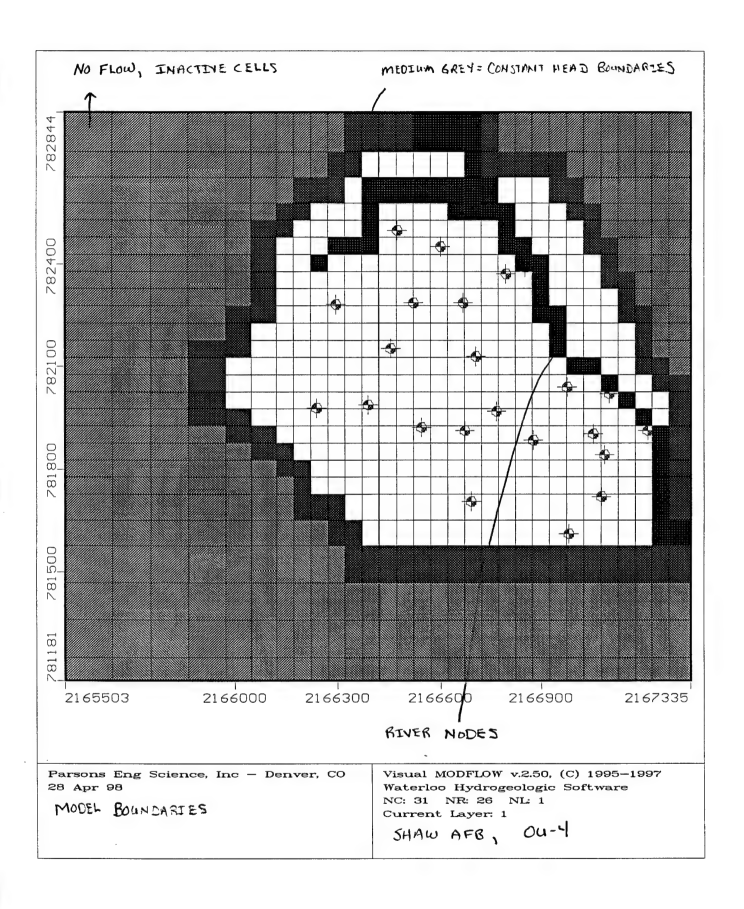
Units for samples are mg/L.
Units for standards are parts per million.

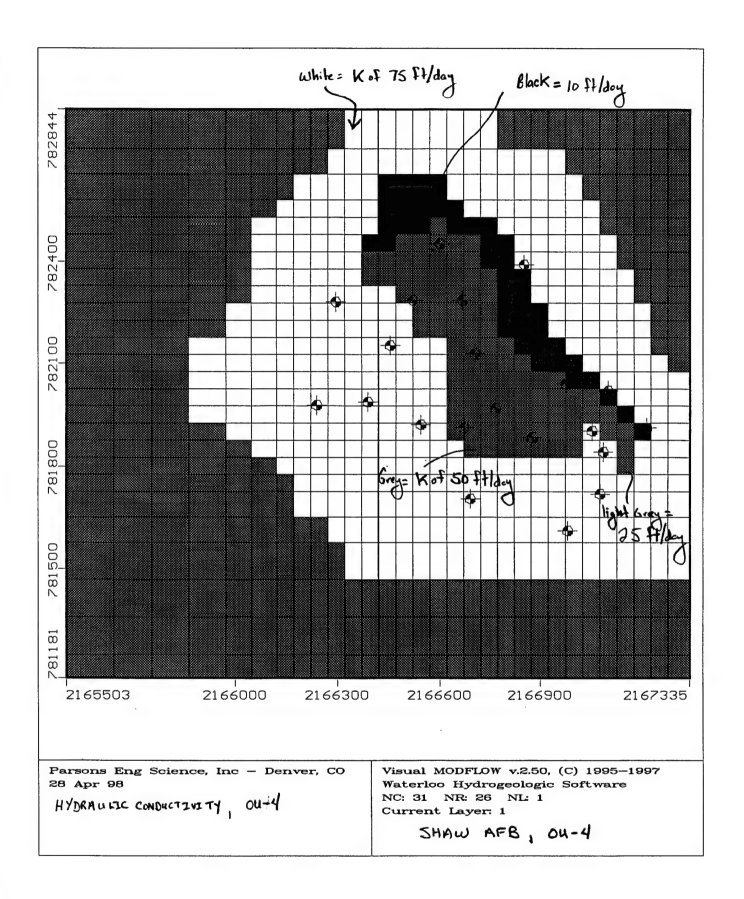
denotes None Detected.

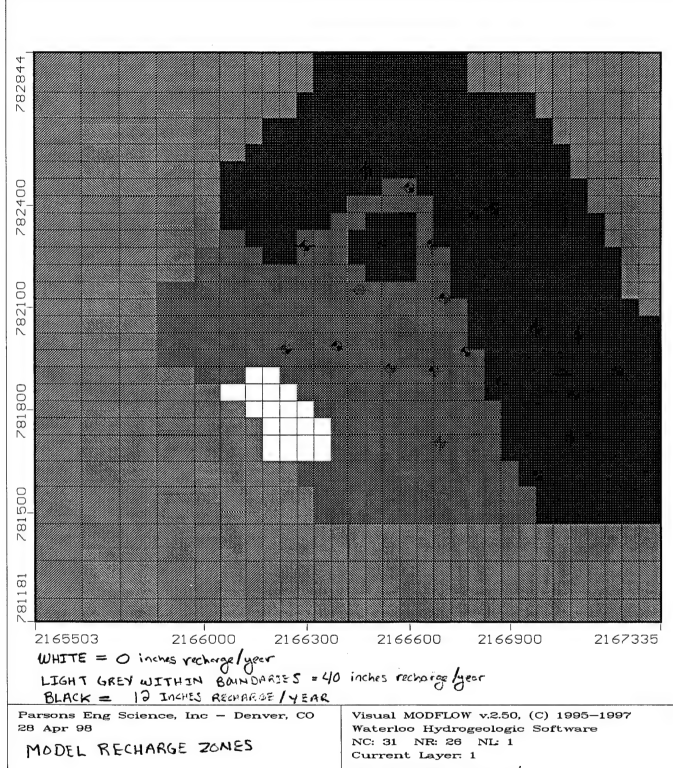
<sup>\*</sup> denotes Below Limit of Quantitation.

# APPENDIX C

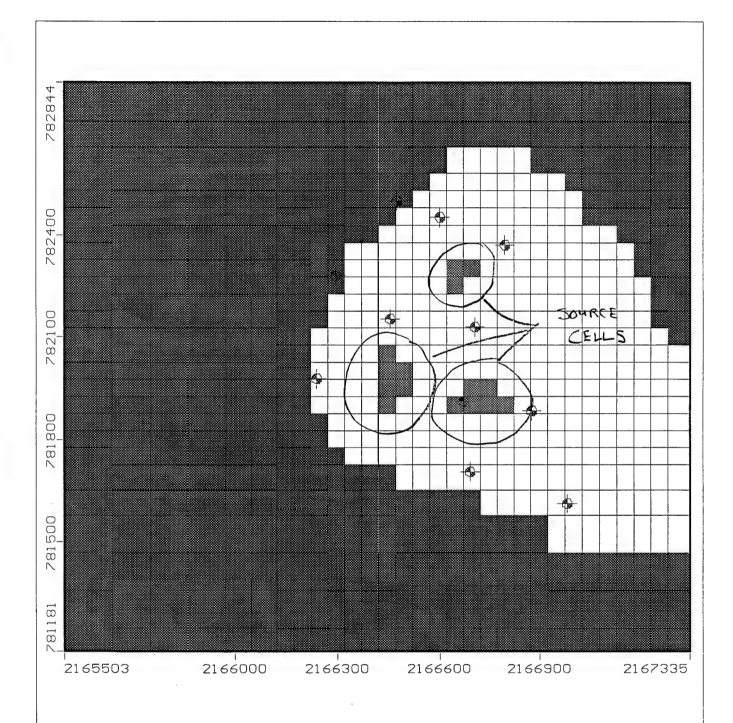
MODEL INPUT PARAMETERS, MODEL OUTPUT AND SUPPORTING FIGURES, AND CALCULATIONS RELATED TO MODEL CALIBRATION.







SHAW AFB, OU-4

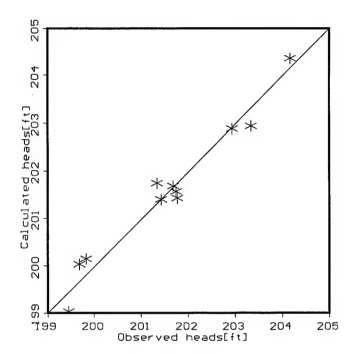


Parsons Eng Science, Inc - Denver, CO 28 Apr 98

SOURCE CELLS FOR CAH
CONTAMINATION

Visual MODFLOW v.2.50, (C) 1995—1997 Waterloo Hydrogeologic Software NC: 31 NR: 26 NL: 1 Current Layer: 1

SHAW AFB, OU-4



Period: 1 Step: 1

Mean error:

-0.0182333

Mean abs. err: 0.244455

RMS error: 0.289165

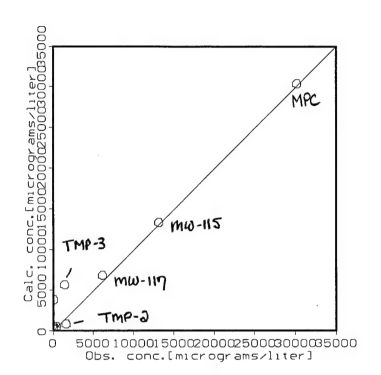
Parsons Eng Science, Inc - Denver, CO 28 Apr 98

CALCULATED VERSUS OBSERVED

HEADS

Visual MODFLOW v.2.50, (C) 1995-1997 Waterloo Hydrogeologic Software NC: 31 NR: 26 NL: 1 Current Layer: 1

SHAW AFB, 04-4



Period: 1 Step: 1

Mean error: 405.45

Mean abs. err: 497.097

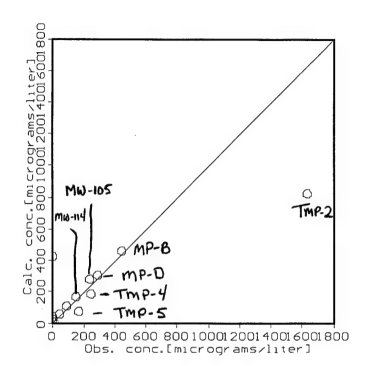
RMS error: 1245.47

Parsons Eng Science, Inc - Denver, CO 28 Apr 98

CALCULATED VERSUS OBSERVED
CAH CONCENTRATIONS
(RANGE O -> 35,000 µg/L)

Visual MODFLOW v.2.50, (C) 1995-1997 Waterloo Hydrogeologic Software NC: 31 NR: 26 NL: 1 Current Layer: 1

SHAW AFB, DU-4



Period: 1 Step: 1

Mean error: 405.45

Mean abs. err: 497.097

RMS error: 1245.47

Parsons Eng Science, Inc - Denver, CO 28 Apr 98

CALCULATED VERSUS OBSERVED

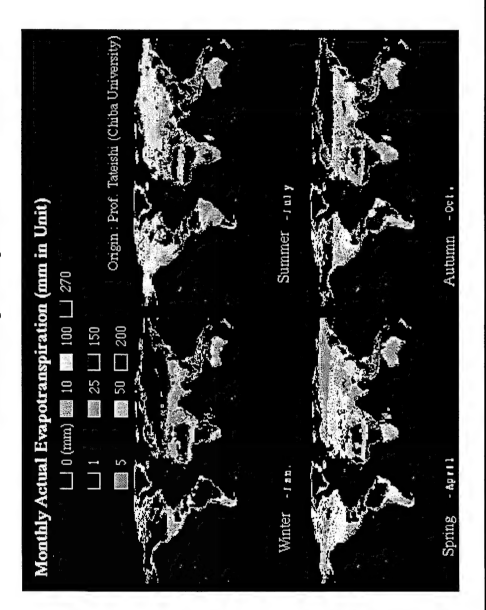
CAH CONCENTRATIONS

Visual MODFLOW v.2.50, (C) 1995-1997 Waterloo Hydrogeologic Software NC: 31 NR: 26 NL: 1 Current Layer: 1

SHAW AFB, OU-4

(RANGE 0 → 1,800 mg/L)

# Potential Evapotranspiration



estimated monthly global potential evapotranspriration(PET), evapotranspiration(ET), water balance on 30-minute latitute-longtitude This data set is produced by AHN and Tateishi(Remote Sensing and Image Research Center, Chiba University, Japan). It consists of atitudinal distribution shows that the larger amount of water surplus in 45N-75N and deficit in 10S-30S rather than those of Legates supplied by GRID-Geneva) data. Almost of all source data were those of approximately 1920 through 1980. The comparison of the Thornthwaite method to estimate evapotranspiration using precipitation (supplied by NOAA-EPA) and soil water holding capacity elevation, which are parts of Global Ecosystems Database supplied by NOAA-EPA. Then further calculation was conducted on grid. Estimation was based on the Priestley-Taylor method using global data sets including air temperature, albedo, cloudiness,

ESTIMATING EVAPOTRASTIRATION RATE TO ESTIMATE A RECHARGE RATE.

(A) SOLVE EVAPOTRASPIRATION RATE TO USE IN HYDRAULIC MASS BALANCE FOR RECHARGE.

ETO (in) = 0.63 [50 (1-32) /9]] x F(2)

Thornthweit's Formula for Estimuty Potential Exapo tenspiration in., de Marsily , G., Orealithtine Hydroges

to Temp in degrees of

I = Amu. I thermal Index = sum of Monthy indexes (i)

Academic Prass, 1986, pp. 403. (Attached)

9= 6.75 x 10-913 -7.71 x 10-513+ 1.70 x 10-01 + 0,49239

F(2) = correction coefficient dependent on latitude: month. Table A.1.1

i= [(+-32)/9]1.5H

Estimbe I

Approxime Linde = 33.8. Al of My Me Bach

	NEAN TEMP.	; **	
Jan.	44.1	1.56	_)
Feb.	47.1	2.19	/
Merch.	22.4	4.25	(
April \	63.1	6.54	
Meg	740	9.21	$\rangle$
June	90.0	11.44	1
July	80.5	13.81	1
Auxus,	77.4	1237	- \
September	74.4	10.45	
October	63.9	6.79	
Nov.	55.4	4.25	- /
Dec.	47.4	2.36	/
VA	e= 63.3°F	(	,

Zi: 84,12 = I

\* Date from Jathard Rajonal Clinck Center in Florence, Sc. 2x See Attented Table Estim-te a

6.75x 10-7 (84,12)3 - 7.71 x 10-5 (84.12)2 + 1.79 x 10-0 (84,12) + 0.45839

0.400 - 0.5456 + 1,506 + 0.41939

1.854

Table to Calculate Monthy Evapotranspiration Rates Myrtle Beach, South Carolina:

Parameters:

84.12 1.854

Month	Mean Temp. (Fahrenheit)	$F(\lambda)$	ETp (inches)
January	44.1	0.88	0.37
February	47.1	0.85	0.53
March	55.4	1.03	1.45
April	63.1	1.09	2.61
May	71	1.2	4.37
June	<b>7</b> 7	1.2	5.70
July	80.5	1.22	6.65
August	79.4	1.16	6.06
September	74.4	1.03	4.38
October	63.9	0.97	2.43
November .	55.4	0.87	1.23
December	47.4	0.86	0.56
		Sum=	36 34

(inches of evapotranspiration per year)

The sec Elect

2 30 16

ظر

The Water Bolance for an annual period is:

P=Q+EV + AS + AS6+R

(in, Groundweter", Freeze : Cherry, 1979, Prantice I-lell, New Joney

P= Praipitation

Forus = D

EY = Evopotranspiration

ASS = change of storage of a surface water reservoir

ASq = Change in storey of a groundwater reservoir

R = Recharge

A) Since we are avanging many years worth of precipitation date, ASS = DSg = O.

B Because the source record is in a depression, little characterists for runoff.
Therefore, Q = 0

( Annual Procipitation is 49.8 inches/you (Section 3)

(Previous Page)

Therefore ...

49.8 = 63 + 36.3 + 485 + 486 + R R = 13.5 inches par year.

13.5/49,8 =0.271 or 27.1% of Amuel Precipitation.

Therefore... it is rewondok to assume that a recharge rate of \$20% por year is possible for two Mystle Barch Sike

Furthermore, on the bosis of Robb C. Heath's paper ["Clossification of Groundwater Systems of the united states", Groundwater, Vol. 30(1), 1982] the astimated aquifor realized rock for the Atlantic Gulf and Coastel Plain ranges from 2-20 inches/your.

Therefore ... 2-2 20 inches/year of recharge is a 4-40% of the annual precipitation for the Mystle Borchance. The above calculated value of 30% is within this range.

50 SHETS FILER SOUNK 1982 100 SHETTS FILE SOUNK 1982 100 SHETTS FILE ASP SOUNK 1982 100 RECYCLED WHITE SOUNK 1995 100 RECYCLED WHI

National Bras

the the control of th

# Thornthwaite's Formula for Estimating Potential Evapotranspiration

## References:

Thornthwaite, C.W., Transactions, Am. Geophys. Union, April 1944, pp. 686-693

Thornthwaite, C.W., The Geographical Review, Vol.38, 1948, pp. 89-94.

de Marsily, G., Quantitative Hydrogeology, Academic Press, 1986, pp. 403-404.

The Thornthwaite formula for monthly potential evapotranspiration, in millimeters per month, is

 $ET_{D}(mm) = 16(10\theta/I)^{a} \times F(\lambda)$ 

Where  $\theta$  is the mean temperature for the month (°C)

 $a = 6.75 \times 10^{-7} I^3 - 7.71 \times 10^{-5} I^2 + 1.79 \times 10^{-2} I + 0.49239$ 

I is an "annual thermal index" equal to the sum of monthly indexes i

 $i = (\theta/5)^{1.514}$ 

 $F(\lambda)$  is a correction coefficient which depends on latitude and month. Values of  $F(\lambda)$  are given in Table A.1.1 of de Marsily (copy included)

For values of temperature, t, in  $^{\circ}F$  and potential evapotranspiration in inches,  $ET_{n}(in)$ , the equations are

$$ET_p(in) = 0.63[50(t-32)/(9I)]^a \times F(\lambda)$$
  
 $i = [(t-32)/9]^{1.514}$ 

Tables 1.1 and 1.2, at the end of this handout, list i for monthly temperatures in °F and °C.

Thornthwaite (1948) states that the equations for potential evapotranspiration given above should only be used for months in which temperatures are 26.5°C (79.7°F) or less. Table 1.3 gives values of ET $_{\rm p}$  for temperatures from 26.5-38.0 °C (79.7-100.4°F).

Table A.1.1. Correction Coefficient  $F(\lambda)$  Depending on the Latitude and the Month

Lat. N.	J	F	M	A	M	J	J	A	S	0	N	D
0	1.04	0.94	1.04	1.01	1.04	1.01	1.04	1.04	1.01			1.04
5	1.02	0.93	1.03	1.02	1.06	1.03	1.06	1.05	1.01			1.02
10	1.00	0.91	1.03	1.03	1.08	1.06	1.08	1.07	1.02			0.99
15	0.97	0.91	1.03	1.04	1.11	1.08	1.12	1.08	1.02			0.97
20	0.95	0.90	1.03	1.05	1.13	1.11	1.14	1.11	1.02			0.94
25	0.93	0.89	1.03	1.06	1.15	1.14	1.17	1.12	1.02			0.91
26	0.92	0.88	1.03	1.06	1.15	1.15	1.17	1.12	1.02	0.99		0.91
27	0.92	0.88	1.03	1.07	1.16	1.15	1.18	1.13	1.02	0.99		0.90
28	0.91	0.88	1.03	1.07	1.16	1.16	1.18	1.13	1.02	0.98	-	0.90
29	0.91	0.87	1.03	1.07	1.17	1.16	1.19	1.13	1.03	0.98		0.89
30	0.90	0.87	1.03	1.08	1.18	1.17	1.20	1.14	1.03	0.98	0.89	0.88
31	0.90	0.87	1.03	1.08	1.18	1.18	1.20	1.14	1.03	0.98	0.89	0.88
32	0.89	0.86	1.03	1.08	1.19	1.19	1.21	1.15	1.03	0.98	0.88	0.87
33	0.88	0.86	1.03	1.09	1.19	1.20	1.22	1.15	1.03	0.97	0.88	0.86
34	0.88	0.85	1.03	1.09	1.20	1.20	1.22	1.16	1.03	0.97	0.87	0.86
35	0.87	0.85	1.03	1.09	1.21	1.21	1.23	1.16	1.03	0.97	0.86	0.85
36	0.87	0.85	1.03	1.10	1.21	1.22	1.24	1.16	1.03	0.97	0.86	0.84
37	0.86	0.84	1.03	1.10	1.22	1.23	1.25	1.17	. 1.03	0.97	0.85	0.83
38	0.85	0.84	1.03	1.10	1.23	1.24	1.25	1.17	1.04	0.96	0.84	0.83
39	0.85	0.84	1.03	1.11	1.23	1.24	1.26	1.18	1.04	0.96	0.84	0.82
40	0.84	0.83	1.03	1.11	1.24	1.25	1.27	1.18	1.04	0.96	0.83	0.81
41	0.83	0.83	1.03	1.11	1.25	1.26	1.27	1.19	1.04	0.96	0.82	0.80
42	0.82	0.83	1.03	1.12	1.26	1.27	1.28	1.19	1.04	0.95	0.82	0.79
43	.0.81	0.82	1.02	1.12	1.26	1.28	1.29	1.20	1.04	0.95	0.81	0.77
44	0.81	0.82	1.02	1.13	1.27	1.29	1.30	1.20	1.04	0.95	0.80	0.76
45	0.80	0.81	1.02	1.13	1.28	1.29	1.31	1.21	1.04	0.94	0.79	0.75
46	0.79	0.81	1.02	1.13	1.29	1.31	1.32	1.22	1.04	0.94	0.79	0.74
47	0.77	0.80	1.02	1.14	1.30	1.32	1.33	1.22	1.04	0.93	0.78	0.73
48	0.76	0.80	1.02	1.14	1.31	1.33	1.34	1.23	1.05	0.93	0.77	0.72
49	0.75		1.02	1.14	1.32	1.34	1.35	1.24	1.05	0.93	0.76	0.71
50	0.74		1.02	1.15	1.33	1.36	1.37	1.25	1.06	0.92	0.76	0.70
Lat. S.												
. 5	1.06	0.95	1.04	1.00	1.02	0.99	1.02	1.03	1.00	1.05	1.03	. 1.06
10	1.08			0.99	1.01	0.96		1.01	1.00	1.06		1.10
15	1.12				0.98				1.00	1.07		1.12
20	1.14				0.96					1.08		1.15
25	1.17				0.94					1.10	1.11	1.18
30	1.17										1.14	1.21
	1.20										1.17	1.25
35	1.23										1.20	1.29
40												1.3
42	1.28											
44	1.30											1.3
46	1.3											1.3
48	1.3					•						
50	1.3	7 1.13	2 1.08	U.0>	0.7	, 0.0		. 5.5				

<sup>&</sup>lt;sup>e</sup> Thornthwaite's formula, from Brochet and Gerbier (1974).

Table 1.1 - Monthly indexes, i, for mean temperature in °C

TOC 0.00 0.10 0.20 0.30 0.40 0.50 0.60 0.70 0.80 0.90 0 0.00 0.01 0.01 0.02 0.03 0.04 0.05 0.06 0.07 1 0.09 0.10 0.12 0.13 0.15 0.16 0.18 0.20 0.23 0.21 2 0.25 0.27 0.29 0.31 0.33 0.35 0.37 0.39 0.42 0.44 3 0.46 0.49 0.51 0.53 0.56 0.58 0.61 0.63 0.66 0.69 4 0.71 0.74 0.77 0.80 0.82 0.85 0.88 0.91 0.94 0.97 5 1.00 1.03 1.06 1.09 1.12 1.16 1.19 1.22 1.25 1.28 б 1.32 1.35 1.39 1.42 1.45 1.49 1.52 1.56 1.59 1.63 7 1.66 1.70 1.74 1.77 1.81 1.85 1.89 1.92 1.96 2.00 8 2.04 2.08 2.11 2.15 2.19 2.23 2.27 2.31 2.35 2.39 9 2.44 2.48 2.52 2.56 2.60 2.64 2.68 2.73 2.77 2.81 10 2.86 2.90 2.94 2.99 3.03 3.08 3.12 3.16 3.21 3.25 11 3.30 3.34 3.58 3.39 3.44 3.48 3.53 3.62 3.67 3.72 12 3.76 3.81 3.86 3.91 3.96 4.00 4.05 4.10 4.20 4.15 13 4.25 4.30 4.35 4.40 4.45 4.50 4.55 4.60 4.65 4.70 14 4.75 4.80 4.86 4.91 4.96 5.07 5.01 5.12 5.17 5.22 15 5.28 5.33 5.38 5.44 5.49 5.55 5.60 5.65 5.71 5.76 16 5.82 5.87 5.93 5.98 6.04 6.10 6.15 6.21 6.26 6.32 17 6.38 6.43 6.49 6.55 6.61 6.66 6.72 6.78 6.84 6.90 18 6.95 7.01 7.07 7.13 7.19 7.25 7.31 7.37 7.43 7.49 19 7.55 7.61 7.67 7.73 7.79 7.85 7.91 7.97 8.03 8.10 20 8.16 8.22 8.28 8.34 8.41 8.47 8.59 8.53 8.66 8.72 21 8.78 8.85 8.91 8.97 9.04 9.10 9.16 9.23 9.29 9.36 22 9.42 9.49 9.55 9.62 9.68 9.75 9.81 9.88 9.95 10.01 23 10.08 10.15 10.21 10.28 10.35 10.41 10.48 10.55 10.61 10.68 10.75 10.82 10.89 10.95 11.02 11.09 11.16 11.23 11.30 11.37 25 11.44 11.50 11.57 11.64 11.71 11.78 11.85 11.92 11.99 12.06 26 12.13 12.21 12.28 12.35 12.42 12.49 12.56 12.63 12.70 12.78 27 12.85 12.92 12.99 13.07 13.14 13.21 13.28 13.36 13.43 13.50 28 13.58 13.65 13.72 13.80 13.87 13.94 14.02 14.09 14.17 14.24 29 14.32 14.39 14.47 14.54 14.62 14.69 14.77 14.84 14.92 14.99 30 15.07 15.15 15.22 15.30 15.38 15.45 15.53 15.61 15.68 15.76 31 15.84 15.91 15.99 16.07 16.15 16.23 16.30 16.38 16.46 16.54 32 16.62 16.70 16.77 16.85 16.93 17.01 17.09 17.17 17.25 17.33 33 17.41 17.49 17.57 17.65 17.73 17.81 17.89 17.97 18.05 18.13 34 18.21 18.30 18.38 18.46 18.54 18.62 18.70 18.79 18.87 18.95 35 19.03 19.11 19.20 19.28 19.36 19.44 19.53 19.61 19.69 19.78 36 19.86 19.94 20.03 20.11 20.20 20.28 20.36 20.45 20.53 20.62 37 20.70 20.79 20.87 20.96 21.04 21.13 21.21 21.30 21.38 21.47 38 21.56 21.64 21.73 21.81 21.90 21.99 22.07 22.16 22.25 22.33 39 22.42 22.51 22.59 22.68 22.77 22.86 22.94 23.03 23.12 23.21 40 23.30 23.38 23.47 23.56 23.65 23.74 23.83 23.92 24.00 24.09

Table 1.2 - Monthly indexes, i, for mean temperature in °F

TOF	0.00	0.20	0.40	0.60	0.80	TOF	0.00	0.02	0.04	0.06	0.08	
32		0.00	0.01	0.02	0.03	66	7.48	7.55	7.61	7.68	7.75	
33	0.04	0.05	0.06	0.07	0.09	67	7.82	7.88	7.95	8.02	8.09	
34	0.10	0.12	0.14	0.15	0.17	68	8.16	8.23	8.29	8.36	8.43	,
35	0.19	0.21	0.23	0.25	0.27	69	8.50	8.57	8.64	8.71	8.78	•
36	0.29	0.32	0.34	0.36	0.39	70	8.85	8.92	8.99	9.07	9.14	
37	0.41	0.44	0.46	0.49	0.51	71	9.21	9.28	9.35	9.42	9.50	
38	0.54	0.57	0.60	0.63	0.65	72	9.57	9.64	9.71	9.79	9.86	
39	0.68	0.71	0.74	0.77	0.81	73	9.93	10.01	10.08		10.23	
40	0.84	0.87	0.90	0.93	0.97	74	10.30			10.52	10.60	
41	1.00	1.03	1.97	1.10	1.14	75				10.90		
42	1.17	1.21	1.24	1.28	1.32	76	11.05	11.13	11.21	11.28	11.36	
43	1.36	1.39	1.43	1.47	1.51	77	11.44	11.51	11.59	11.67	11.74	
44	1.55	1.59	1.62	1.66	1.70	78	11.82	11.90	11.98	12.06	12.13	
45	1.75	1.79	1.83	1.87	1.91	79	12.21	12.29	12.37	12.45	12.53	
46	1.95	1.99	2.04	2.08	2.12	80	12.61	12.69	12.77	12.85	12.93	
47	2.17	2.21	2.26	2.30	2.34	81	13.01	13.09	13.17	13.25	13.33	
48		2.44	2.48	2.53	2.57	82	13.41	13.49	13.58	13.66	13.74	
49		2.67	2.71	2.76	2.81	83	13.82	13.90	13.99	14.07	14.15	
50	2.86	2.90	2.95	3.00	3.05	84	14.23	14.32	14.40	14.48	14.57	
51	3.10	3.15	3.20	3.25	3.30	85	14.65	14.73	14.82	14.90	14.99	
52	3.35	3.40	3.45	3.50	3.55	86	15.07	15.15	15.24	-		
53	3.61	3.66	3.71	3.76	3.82	87			15.67		15.84	
54	3.87	3.92	3.98	4.03	4.09	88	15.92			16.18		
55	4.14	4.19	4.25	4.30	4.36	89	16.36	16.44	16.53		16.70	
56	4.41	4.47	4.53	4.58	4.64	90			16.97		17.14	
57	4.70	4.75	4.81	4.87	4.93	91	17.23		17.41		17.59	
58	4.98	5.04	5.10	5.16	5.22	92		17.77			18.03	
59	5.28	5.34	5.40	5.46	5.52	93		18.21			18.49	
60	5.58	5.64	5.70	5.76	5.82	94		18.67				
61	5.88	5.94	6.00	6.06	6.13	95		19.12			19.40	
62	6.19	6.25	6.31	6.38	6.44	96	19.49	19.58			19.86	
63	6.50	6.57	6.63	6.70	6.76	97	19.95			20.23		
64	6.82	6.89	6.95	7.02	7.08	98		20.51			20.80	
65	7.15	7.22	7.28	7.35	7.41	99	20.89	20.99	21.08	21.17	21.27	

Table 1.3 - Potential evapotranspiration for temperatures greater than 26.5  $^{\circ}\text{C}$ 

T(°C)	ET (mm)	T(°F)	ET (in)
26.5	135.0	79.7	5.315
27.0	139.5	80.6	5.492
27.5	143.7	81.5	5.657
28.0	147.8	82.4	5.819
28.5	151.7	83.3	5.972
29.0	155.4	84.2	6.118
29.5	158.9	85.1	6.256
30.0	162.1	86.0	6.382
30.5	165.2	86.9	6.504
31.0	168.0	87.8	6.614
31.5	170.7	88.7	6.720
32.0	173.1	89.6	6.815
32.5	175.3	90.5	6.902
33.0	177.2	91.4	6.976
33.5	179.0	92.3	7.047
34.0	180.5	93.2	7.106
34.5	181.8	94.1	7.157
35.0	182.9	95.0	7.201
35.5	183.7	95.9	7.232
36.0	184.3	96.8	7.256
36.5	184.7	97.7	7.272
37.0	184.9	98.6	7.280
37.5	185.0	99.5	7.283
38.0	185.0	. 100.4	7.283

# Blaney and Criddle Formula for Consumptive Use (Evapotranspiration)

## Reference:

Blaney, H. 7. and W. D. Criddle, Determining water requirements in irrigated areas from climatological and irrigation data, Soil Conservation Service Technical Paper 96, revised 1952, pp. 14-18.

The empirical formulas developed by Thornthwaite were based on data for relatively humid areas. For arid areas, requiring irrigation for agriculture, Blaney and Criddle developed an alternative formula that relates potential evapotranspiration to the type of crop planted. Their basic formula is as follows:

u = kf

where u is the monthly consumptive use (evapotranspiration) in inches

 $f = (t \times p)/100$  the monthly consumptive use factor

t = mean monthly temperature, °F

p = monthly percent of daytime hours of the year

k = monthly consumptive use coefficient

A graph of monthly percent of daytime hours of the year is included as well as a table (labeled Table 3) of consumptive use coefficients for various crops, averaged over the growing season. Consumptive use is presumably minimal except during the growing season.

### Soil Moisture Retention

The table labeled Table 3 gives estimates of the soil moisture retained in the root zone (soil moisture storage capacity) for various soil types and vegetation types. Note that soil moisture retention increases with decreasing grain size, from 1.2 in/ft for sandy soil to 3.6 in/ft for clayey soil, and with increasing depth of the root zone.

Tuble 3.--Consumptive-use coefficients (K) for irrigated crops in western States

	1.	
Сгор	Length of growing season or period	Consumptive-use coefficient <sup>1</sup> (K)
	Between frosts	0.80 to 0.85
Alfalfa	Between 110313	
Alfalfa.	Prefrost period	-60
Beans	3 months	.60 to .70
Corn	4 months	.75 to .85
Cotton	7 months	.60 to .65
Flax	7 to 8 months	.80
Grains, small	3 months	.75 to .85
Grain (Spring)	3 months	.70
Grain sorghums	4 to 5 months	.70
Orchard, citrus	7 months	.50 to .65
Orchard, walnuts .	Between frosts	.70
Orchard, deciduous	Between frosts	.60 to .70
Pasture, grass	Between frosts	75
Pasture, Ladino clover	Between frosts	.80 to .85
Peas	2 months	.80
Potatoes	3½ months	.65 to .75
Rice	3 to 5 months	.85 to 1:20
Sugar beets	6 months	.65 to .75
Sugarcane (Puerto Rico)	Annual	.80
Tomatoes	4 months	.70
Truck - small	3 months	.60

Average for the growing season or period. Consumptive-use coefficients vary throughout the growing season (see Table 8). The lower values of K are for coastal areas, the higher values for areas with an arid climate.

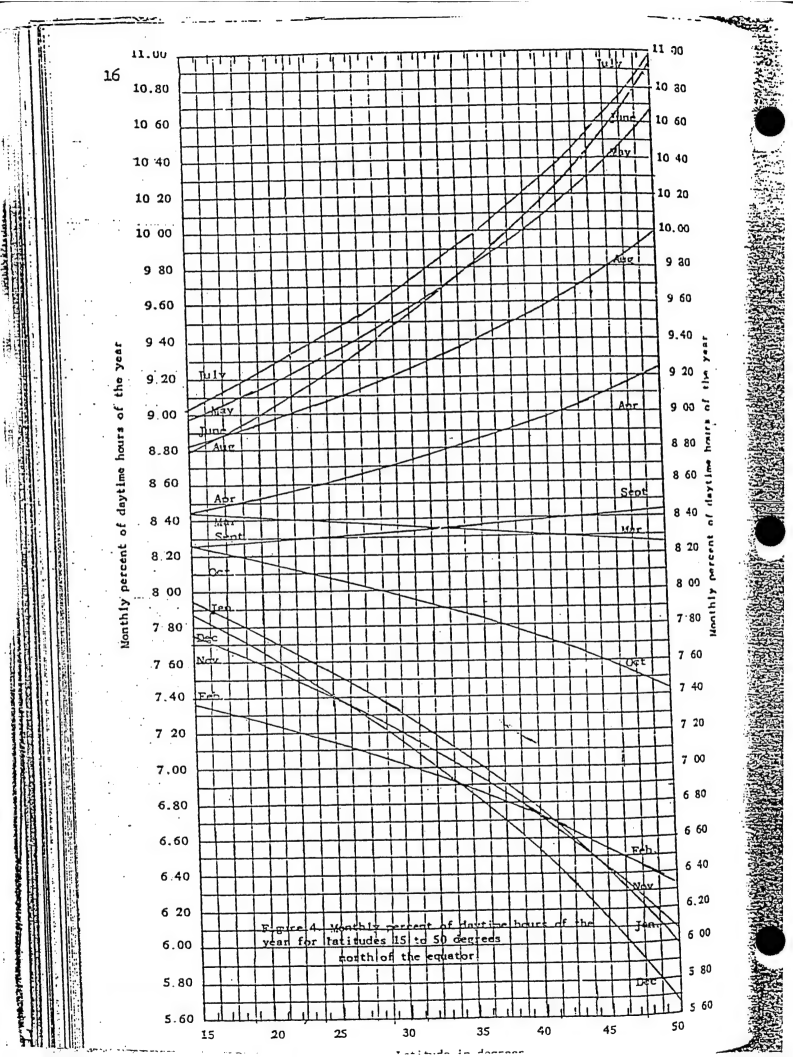


TABLE 10

PROVISIONAL WATER HOLDING CAPACITIES WITH DIFFERENT COMBINATIONS
OF SOIL AND VEGETATION

Once Tring		- 14	Page 1	Jawa	APPLICABLE SOLI	
SOIL TYPE	AVAILABL	-	- Root	-	RETENTIO	
	nn/n	IN/FT	H	FT	HH	( X
SHALLOW-ROOTED CROPS	(SPINACH.	PEAS. BEAR	IS, BEETS, CARRO	T3, ETC.)		
FINE BAND	100	1.2	.50	1.67	50	2.0
FINE SANDY LOAM	150	1.8	.50	1.67	75	3.0
SILT LOAM	200	2.4	. 62	2.08	125	5.0
CLAY LOAM	250	3.0	.40	1.33	100	4.0
CLAY	300	3. ô	.25	.83	75	3.0
MODERATELY DEEP-ROOT	ED CROPS (C	ORM. COTTO	DH. TOBACCO, CER	EAL GRAIRS	)	
FIRE SAND	100	1.2	.75	2,50	75	3.0
FINE SANDY LOAM	150	1.8	1.00	3.33	150	6.0
SILT LOAH	200	2.4	1.00	3.33	200	0.3
CLAY LOAH	250	3_0	.80	2.67	200	8.0
CLAY	300	3.6	•50	1.67	150	6.0
DEEP-ROOTED CROPS (A	LFALFA, PA	STURES, SH	RUBS)			
FERE SAND	100	1.2	1.00	3.33	. 100	4.0
FINE SANDY LOAM	150	1.8	1.00	3.33	150	6.0
SILT LOAM	200	2.4	1.25	4.17	250	10.0
CLAY LOAM	250	3.0	1.00	3.33	250	10.0
CLAY .	300	3.6	. 67	2.22	200	8.0
ORCHARDS						
FINE SAND	100	1.2	1.50	5.00	150	6.0
FINE SANDY LOAM	150	1.8	1.67	5.55	250	10.0
SILT LOAM	200	2.4	1.50	5.00	300	12.0
CLAY LOAH	250	3.0	. 1.00	3, 33	250	10.0
CLAY	300	3.6	.67	2,22	200	8.0
CLOSED HATURE FORES	r					
FIRE SAND	100	1.2	2,50	8, 33	250	10.0
FIRE SAKOY LOAM	150	1.8	2.00	. 6, 66	300	12.0
SILT LOAM	200	2.4	2.00	6.65	400	16.0
CLAY LOAM	250	3.0	1.60	5.33	400	163
CLAY	300	3.6	1.17	3.90	350	14.0

THESE FIGURES ARE FOR HATURE VEGETATION. YOUNG CULTIVATED CROPS, SEEDLINGS, AND OTHER IMMATURE VEGETATION WILL HAVE SHALLOWER ROOT ZONES AND, HENCE, MAYE LESS WATER AVAILABLE FOR THE USE OF THE VEGETATION. AS THE PLANT DEVELOPS FROM A SEED OR A YOUNG SPROUT TO THE MATURE FORM, THE ROOT ZONE WILL INCREASE PROGRESSIVELY FROM ONLY A FEW INCHES TO THE VALUES LISTED ABOVE. USE OF A SERIES OF SOIL MOISTURE RETERTION TABLES WITH SUCCESSIVELY INCREASING VALUES OF AVAILABLE MOISTURE PERMITS THE SOIL MOISTURE TO BE DETERMINED THROUGHOUT THE GROWING SEASON.

# SUMMARY OF AVERAGE AN EIGHTED AVERAGE PERCENTAGES OF FOR THE TOTAL CHLORINATED PLUME (INCLUDES ETHENES AND ETHANES) SITE OU-4 RNA TS

						ZHZ	SHAW AFB, SOUTH CAROLINA	UTH CARO	LINA						
(¥	<u>@</u>	(0)	<u>(a</u> )	(E)	(F)	9	Ξ	€	3	( <u>X</u>	(-)	(W)	(X)	<u>(</u> 0	(A)
	Total	PCE/	Column C	TCE/	Column E	Cis-1,2/	Column G	1,1-DCE/	Column I	/C	Column K	<b>1</b> C₹	Column M	1,1-DCA/	Column O
Sample	CAH	Total	×	Total		Total		Total		Total		Total		Total	
· A	(μg/L)	CAH	Column B	CAH	Column B	CAH	Column B	CAH	Column B	CAH	Column B	CAH	Column B	CAH	Column B
MW-105	236.5	0.42%	1.00	13.23%	31.30	84.99%	201.00	0.42%	1.00	0.51%	1.20		•	0.45%	1.00
MW-106	9.6		,			29.60%	5.90	10.10%	1.00					30.30%	3.00
MW-107	2.6		•			27.69%	1.50		•				,	42.31%	1.10
MW-111	-	•	•	100.00%	1.00	•	1						•	•	,
MW-112	06		•		•	24.44%	22.00	2.44%	2.20			1.11%	1.00	72.00%	64.80
MW-112A	0	1		٠	•	,	٠	,	1	•		,		•	
MW-113	2.2	1	•	•		54.55%	1.20	,					•	45.45%	1.00
MW-114	148		•		,	58.85%	87.10	1.42%	2.10				•	39.73%	58.80
MW-115	13149	0.04%	4.80	0.08%	10.20	12.32%	1620.00	2.80%	368.00	3.16%	416.00	15.44%	2030.00	66.16%	8700.00
MW-115A	4.1		,	24.39%	1.00			51.22%	2.10	1			,	24.39%	1.00
MW-116	0		,			•	,	,					,	,	
MW-116A	0				•		•	•	•			•			
MW-117	6087.7	0.40%	24.10	3.33%	203.00	7.51%	457.00	2.43%	148.00	1.39%	84.60	77.04%	4690.00	7.90%	481.00
MW-118	-		1	•	•		•	•	•			100.00%	1.00	•	
TW-101	o		. 1				•	•	•	,				•	
TW-120	0	•	•		•		•		•					•	
TW-121	56.9	1				53.95%	30.70	12.83%	7.30	26.89%	15.30		•	2.64%	1.50
MPA	48		•	2.08%	1.00	7.29%	3.50	8.96%	4.30			17.50%	8.40	64.17%	30.80
MPB	444	0.23%	1.00	0.29%	1.30	2.39%	10.60	1.06%	4.70	,		92.57%	411.00	3.47%	15.40
MPC	30237.1	0.30%	90.70	2.37%	718.00	15.18%	4590.00	2.47%	748.00	0.10%	30.40	76.07%	23000.00	3.51%	1060.00
MPD	288.3	•	•	26.71%	77.00	53.76%	155.00	0.59%	1.70			0.45%	1.30	18.49%	53.30
MPE	7.3	•	•		•	67.12%	4.90	•	•			13.70%	1.00	19.18%	1.40
TMP-1	0	•	,	•	•	,	•	,	,				•	,	
TMP-2	1628.6	0.06%	1.00	0.07%	1.20	20.82%	339.00	7.12%	116.00	1.40%	22.80	4.83%	78.60	65.70%	1070.00
TMP-3	1468.2	0.07%	1.00	0.27%	3.90	10.22%	150.00	2.74%	40.20	4.06%	29.60	4.12%	60.50	78.53%	1153.00
TMP-4	247.1			0.40%	1.00	68.39%	169.00	1.42%	3.50	3.12%	7.70	0.40%	1.00	26.26%	64.90
TMP-5	163.8	•	٠			25.95%	42.50	1.10%	1.80	70.82%	116.00			2.14%	3.50
	Average:	0.22%		14.44%		38.06%		6.82%		12.38%		33.60%		32.25%	
	Weighted Average:	verage:	0.23%		1.93%		14.53%		2.67%		1.39%		55.75%		23.50%
Migration.	Pathway Alon,	g East/South	Migration Pathway Along East/Southeast Migration Lobe	rope											
MPC	30237.1		90.70	2.37%	718.00	15.18%	4590.00	2.47%	748.00	0.10%	30.40	76.07%	23000.00	3.51%	1060.00
MW-115	13149	0.04%	0.80	0.08%	10.20	12.32%	1620.00	2.80%	368.00	3.16%	416.00 59.60	15.44%	2030.00	66.16%	8700.00
IMF-3	1400.2		5.	0.4170	3.30	10.62 /0	100.00	6.1.1.2 0.000	40.40	4.00 %	00.00	21 070	00.00	10.00 /0 10.40 /0	100.00
	Average:	0.13%	0 22	0.91%	1 63 0	12.57%	14 100	7.01%	2000	7.44%	1120	31.87%	25.040	49.40%	24 22 00
	Weighted Average:	verage:	0.22%		1.63%		14.18%		2.58%		1.13%		55.94%		24.33%

# GROUNDWATER QUALITY DATA SUMMARY FOR CHLORINATED ETHENES AND ETHANES SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

10131	250	(μg/L)	236.5	6.6	2.6	-	90	0	2.2	148	13149	4.1	0	0	6087.7	-	0	0	56.9	48	444	30237.1	288.3	7.3	0	1628.6	1468.2	247.1	163.8			
-1,1-	e C	(µg/L)	1	3.0	1.1	Q	64.8	Q	1	58.8	8700	-	S	£	481	S	S	Ę	1.5	30.8	15.4	1060	53.3	1.4	Ð	1070	1153	64.9	3.5	1060	8700	1153
1,1,1	ICA V	(μg/L)	QN	S	S	S	-	Ð	£	S	2030	S	S	S	4690	1	2	S	Q.	8.4	411	23000	1.3	1	Q	78.6	60.5	-	QN	23000	2030	60.5
Vinyl	Chloride	(μg/L)"	1.2	S	2	2	Q.	Q.	Ð	g	416	Q	S	Q	84.6	Ð	Q	Q	15.3	S	S	30.4	2	S	S	22.8	59.6	7.7	116	30.4	416	59.6
-1,1-	E C	(μg/L)	1	-	£	S	2.2	2	S	2.1	368	2.1	Ş	£	148	Ð	£	S	7.3	4.3	4.7	748	1.7	Ð	Ð	116	40.2	3.5	1.8	748	368	40.2
trans-	1,2-DCE	(µg/L)	ND.	Q.	Q.	Q.	S	S	Ş	Š	Ą	£	£	<del>S</del>	Ę	£	Ð	£	2.1	Ð	S	S S	£	Ę	ğ	S S	Ž	Ę	ND	Q	QX	2
CIS-	1,2-DCE	(μg/L)	201	5.9	1.5	N S	22.0	N Q	1.2	87.1	1620	Q	R	Ę	457	S S	£	S	30.7	3.5	10.6	4590	155	4.9	N <sub>D</sub>	339	150	169	42.5	4590	1620	150
	E C E	(μg/L)	31.3	Q	R	_	S	2	S	S	10.2	1	Q.	QN	203	S	Q	Q.	QN.	-	1.3	718	77.0	g	Q	1.2	3.9	-	QN	718	10.2	3.9
Š	FCE.	(µg/L)	-	£	Q.	R	Q.	Z	Z	R	8.4	R	S	£	24.1	£	S	Ð	Ð	£	-	7.06	Ð	Ð	R	1	1.0	Q.	æ	90.7	8 4	1.0
	Date	Sampled	5/15/97	5/14/97	5/14/97	5/15/97	5/16/97	5/16/97	5/16/97	5/15/97	5/15/97	5/15/97	5/14/97	5/14/97	5/15/97	5/15/97	5/14/97	5/17/97	5/17/97	5/16/97	5/16/97	5/16/97	5/16/97	5/17/97	5/17/97	2/11/67	2/11/97	2/16/97	2/16/97	2/16/97	2/15/97	5/11/97
	Sample	О	MW-105	MW-106	MW-107	MW-111	MW-112	MW-112A	MW-113	MW-114	MW-115	MW-115A	MW-116	MW-116A	MW-117	MW-118	TW-101	TW-120	TW-121	MPA	MPB	MPC	MPD	MPE	TMP-1	TMP-2	TMP-3	TMP-4	TMP-5	MPC	MW-115	TMP-3

# SUMMARY OF AVERAGE AND WHITED AVERAGE PERCENTAGES OF INDIVIDUAL CHLORINATED ETHENE AND ETHANE PLUMES SITE OU 4 RNA TS

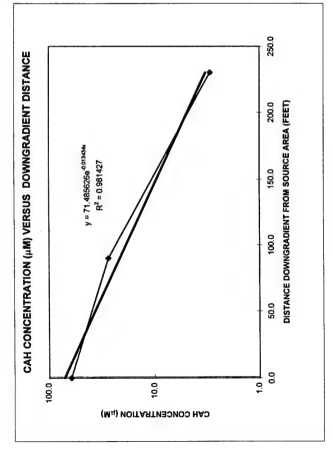
							SHAW	SHAW AFB, SOUTH CAROLINA	CAROLINA							
8	(B)	(0)	0	<u>(i)</u>	(F)	(0)	Ξ	(3)	3	<del>(X</del>	(3)	(W)	(S	(0)	(P)	(O)
	Total	Total	PCE/		TCE/		Cis-1,2/		1,1-DCE/		/C		TCA		1,1-DCA/	
Sample	Ethenes	Ethanes	Total	Column D	Total	Column F		Column H	Total	Column J	Total	Column L	Total	Column N	Total	Column P
a	(μg/L)	(μg/L)	Ethene	x Column B	Emene	x Column B	-	x Column B	Emene	x Column B	Emene	x Column B	Ethane	x column x	Emane	y column y
MW-105	235.5		0.42%	1.00	13.29%	31.30	85.35%	201.00	0.42%	00.1	%LC:0	02.1			100.00%	9.
MW-106	6.9	3	•	•			85.51%	5.90	14.49%	1.00		•			100.00%	3.00
MW-107	1.5	Ξ	,		•		100.00%	1.50	•		•		•	,	100.00%	1.10
MW-111	-	0			100.00%	1.00		•		•	•	•	•			•
MW-112	24.2	65.8	•		•		90.91%	22.00	%60.6	2.20	•	,	1.52%	1.00	98.48%	64.80
MW-112A	0	0	,				,		•		•		•	•		
MW-113	1.2	-	,				100.00%	1.20	•		•			•	100.00%	1.00
MW-114	89.2	58.8		•			97.65%	87.10	2.35%	2.10	•			•	100.00%	58.80
MW-115	2419	10730	0.20%	4.80	0.42%	10.20	66.97%	1620.00	15.21%	368.00	17.20%	416.00	18.92%	2030.00	81.08%	8700.00
MW-115A	3.1	1		•	32.26%	1.00	•		67.74%	2.10				•	100.00%	1.00
MW-116	0	0			•	•			•		•					
MW-116A	0	0			•	•	•				•			•		•
MW-117	916.7	5171	2.63%	24.10	22.14%	203.00	49.85%	457.00	16.14%	148.00	9.23%	84.60	90.70%	4690.00	9.30%	481.00
MW-118	0	1		•				•			•		100.00%	1.00		•
TW-101	0	0		•					,		,					,
TW-120	0	0	•	•					•		•			•		
TW-121	55.4	1.5	,	•		•	55.42%	30.70	13.18%	7.30	27.62%	15.30	•		100.00%	1.50
MPA	80.00	39.2	,	•	11.36%	1.00	39.77%	3.50	48.86%	4.30		,	21.43%	8.40	78.57%	30.80
MPB	17.6	426.4	2.68%	1.00	7.39%	1.30	60.23%	10.60	26.70%	4.70	•		96.39%	411.00	3.61%	15.40
MPC	6177.1	24060	1.47%	90.70	11.62%	718.00	74.31%	4590.00	12.11%	748.00	0.49%	30.40	95.59%	23000.00	4.41%	1060.00
MPD	233.7	54.6	,	•	32.95%	77.00	66.32%	155.00	0.73%	1.70	,		2.38%	1.30	97.62%	53.30
MPE	4.9	2.4				•	100.00%	4.90			1		41.67%	1.00	58.33%	1.40
TMP-1	0	0	•	,			,		,	•	•		,			
TMP-2	480	1148.6	0.21%	1.00	0.25%	1.20	70.63%	339.00	24.17%	116.00	4.75%	22.80	6.84%	78.60	93.16%	1070.00
TMP-3	254.7	1213.5	0.39%	1.00	1.53%	3.90	58.89%	150.00	15.78%	40.20	23.40%	29.60	4.99%	60.50	95.01%	1153.00
TMP-4	181.2	62.9	,	•	0.55%	1.00	93.27%	169.00	1.93%	3.50	4.25%	7.70	1.52%	1.00	98.48%	64.90
TMP-5	160.3	3.5	,	•		•	26.51%	42.50	1.12%	1.80	72.36%	116.00	•	•	100.00%	3.50
		Average:	1.57%		19.48%		73.42%		16.88%		17.76%		40.16%	•	79.90%	
		Weighted Average:	rage:	1.10%		9.31%		70.00%		12.88%		%69.9		70.35%		29.62%
Migration Pathy	way Along East/	Migration Pathway Along East/Southeast Migration Lobe	tion Lobe													
MPC	6177.1	24060	1.47%	90.70	11.62%	718.00	74.31%	4590.00	12.11%	748.00	0.49%	30.40	95.59%	23000.00	4.41%	1060.00
MW-115	2419	10730	0.20%	4.80	0.42%	10.20	%26.99	1620.00	15.21%	368.00	17.20%	416.00	18.92%	2030.00	81.08%	8700.00
TMP-3	254.7	1213.5	0.39%	1.00	1.53%	3.90	58.89%	150.00	15.78%	40.20	23.40%	59.60	4.99%	60.50	95.01%	1153.00
		Average:	%69.0		4.53%		66.72%		14.37%		13.70%		39.83%		60.17%	
		Weighted Average:	rage:	1.09%		8.27%		71.86%		13.06%		5.72%		%69.69		30.31%

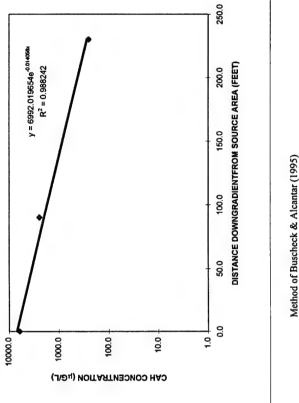
# ESTIMATED MINIMUM FIRST-ORDER RATE CONSTANT CALCULATION FOR THE CHLORINATED ETHENE CONTAMINATION (Foc = 0.00024) USING THE METHOD OF BUSCHECK AND ALCANTAR (1995)

MAY, 1997 SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

	Distance	PCE (µg/L)	PCE (µM)	ΓCE (μg/L	TCE (µM)	TCE (μM) Total DCE (μg/L)	ΛL) Total DCE (μM) VC	VC (µg/L)	VC (µM)	Total CAH (μg/L) To	#
Well	Downgradient	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97
MPC	0.0	90.7	9.0	718.0	5.5	5338.0	55.1	30.4	9.0	6177.1	9.19
MW-115	0.06	8.4	0.0	10.2	0.1	1988.0	20.5	416.0	6.7	2419.0	27.3
TMP-3	230.0	1.0	0.0	3.9	0.0	190.2	2.0	9.69	1.0	254.7	3.0

CAH CONCENTRATION (µG/L) VERSUS DOWNGRADIENT DISTANCE





Method of Buscheck & Alcantar (1995)  $\lambda = v_c/4\alpha_x([1+2\alpha_x(k/v_x)]^2-1)$  where  $v_c = 2.97$  ft/day\*

nete  $V_c = 2.97$  to  $\alpha_x = 50$  ft k/v = -0.01343

therefore  $\lambda = -1.3103E-02$  1/day  $\lambda = -1.3103E+00$  %/day

\*(assumes gw velocity of 3.3 and retardation coefficient of 1.11)

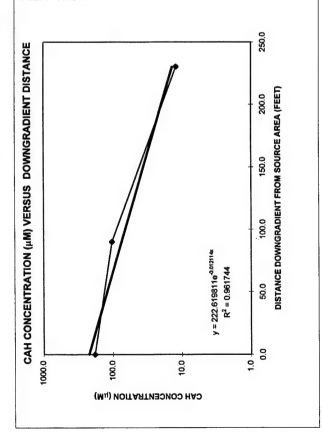
where  $v_c = 2.97$  (f/day\*  $\alpha_x = 2.97$  (f/day\*  $\alpha_x = 5.0$  ff k/v = -0.01405therefore  $\lambda = -1.2414E+00$  %/day

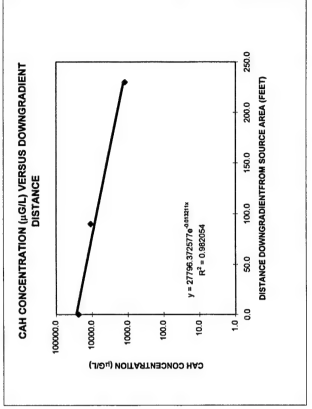
\*(assumes gw velocity of 3.3 and retardation coefficient of 1.11)

# ESTIMATED MINIMUM FIRST-ORDER RATE CONSTANT CALCULATION FOR THE CHLORINATED ETHANE CONTAMINATION (Foc = 0.00024) USING THE METHOD OF BUSCHECK AND ALCANTAR (1995)

MAY, 1997 SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

Well	Distance Downgradient	1,1,1-TCA (µg/L) May-97	1,1,1-TCA (μM) May-97	1,1-DCA (μg/L) May-97	1,1-DCA (μM) May-97	Total CAH (μg/L) May-97	Total CAH (μΜ) May-97
MPC	0.0	23000.0	172.4	1060.0	10.7	24060.0	183.1
MW-115	0.06	2030.0	15.2	8700.0	87.9	10730.0	103.1
rMP-3	230.0	60.5	0.5	1153.0	11.7	1213.5	12.1







where 
$$v_c = 2.92$$
 ft/day\* 
$$\alpha_x = 50$$
 ft 
$$kv = -0.012114$$

therefore 
$$\lambda = -1.3948E-02$$
 1/day  $\lambda = -1.3948E+00$  %/day

\*(assumes gw velocity of 3.3 and retardation coefficient of 1.13)

Method of Buscheck & Alcantar (1995)  $\lambda = v_o/4\alpha_x([1+2\alpha_x(k/v_x)]^2-1)$ 

where 
$$v_c = 2.92$$
 ft/day  $\alpha_x = 50$  ft  $kv = -0.013211$ 

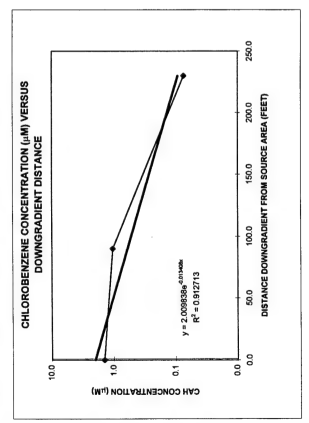
therefore 
$$\lambda = -1.3095E-02$$
 1/day  $\lambda = -1.3095E+00$  %/day

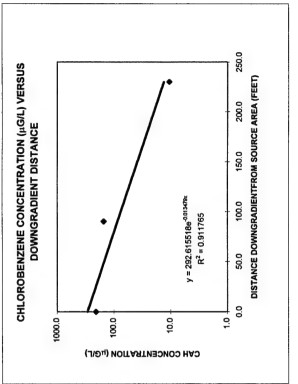
\*(assumes gw velocity of 3.3 and retardation coefficient of 1.13)

# ESTIMATED MINIMUM FIRST-ORDER RATE CONSTANT CALCULATION FOR THE CHLORINATED BENZENE CONTAMINATION (Foc = 0.00024) USING THE METHOD OF BUSCHECK AND ALCANTAR (1995) MAY, 1997

# MAY, 1997 SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

	Distance	1,2-DCB (µg/L) 1,2-DCB		1,3-DCB (µg/L)	1,3-DCB (µM)	1,4-DCB (µg/L)	1,4-DCB(μM) CB (μg/L CB (μM) Total C	CB (µg/L	CB (µM)	Total C-Benz. (µg/L)	Total C-Benz. (µM)
Well	Downgradient	May-97	May-97	May-97	May-97	May-97	May-97	May-97 May-97	May-97	May-97	May-97
MPC	0.0	117.0	8.0	14.1	0.1	68.2	9.5	9.2	0.1	208.5	1.4
MW-115	0.06	88.9	9.0	11.6	0.1	46.4	0.3	4.9	0.0	151.8	1.0
TMP-3	230.0	6.3	0.0	0.0	0.0	3.3	0.0	1.0	0.0	10.6	0.1





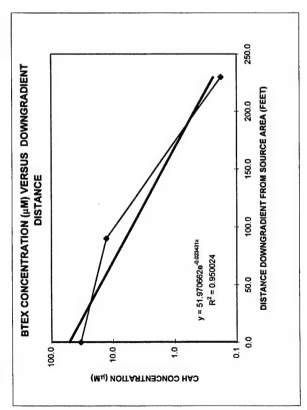
p 25	Method of Buscheck & Alcantar (1995) $\lambda = v_c/4\alpha_x([1+2\alpha_x(k/v_x)]^2-1)$	where $v_c = 2.56$ ft/day* $\alpha_x = 50$ ft k/v = -0.013408	therefore $\lambda = -1.1313E-02$ 1/day $\lambda = -1.1313E+00$ %/day
------	---------------------------------------------------------------------------------------------	---------------------------------------------------------------	-----------------------------------------------------------------------

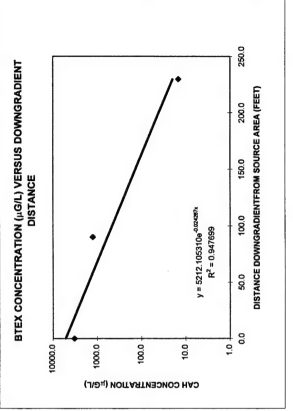
							ent of 1.285)
ar (1995)	J-1)	ft/day	ff		1/day	%/day	dation coefficie
Method of Buscheck & Alcantar (1995)	$\lambda = v_c/4\alpha_x([1+2\alpha_x(k/v_x)]^2-1)$	2.56	20	-0.013479	-1.1251E-02	-1.1251E+00	*(assumes gw velocity of 3.3 and retardation coefficient of 1.285)
hod of B	ک ۱۱ د	ار د	$\alpha_{x} =$	k/v =	<del>ا</del> ا	۳	umes gw v
Met		where $v_c =$			therefore		*(ass

# ESTIMATED MINIMUM FIRST-ORDER RATE CONSTANT CALCULATION FOR THE BTEX CONTAMINATION (Foc = 0.00024) USING THE METHOD OF BUSCHECK AND ALCANTAR (1995) MAY, 1997

SITE OU-4 RNA TS
SHAW AFB, SOUTH CAROLINA

	Distance	Benzene (µg/L)	Benzene (µM)	Toluene (µg/L	uene (µ]	Sth. Benz. (µg/L)	Eth. Benz. (µM)	Xylenes (μg/L	Xylenes (μM)	Xylenes (μM) Total BTEX (μg/L	Total BTEX. (µM)
Well	<b>Downgradient</b>	May-97	May-97	May-97	May-97	May-97	May-97 May-97 May-97	May-97	May-97	May-97	May-97
MPC	0.0	81.7	1.0	1685.0	18.3	242.0	2.3	1279.0	12.0	3287.7	33.7
MW-115	90.0	157.0	2.0	393.0	4.3	113.0	1.1	588.0	5.5	1251.0	12.9
TMP-3	230.0	12.4	0.2	0.0	0.0	0.0	0.0	2.2	0.0	14.6	0.2







ft/day* ft	herefore $\lambda = 1.3979E-02$ 1/day $\lambda = 1.3979E+00$ %/day
2.7 50 -0.024267	1.3979E-02 1.3979E+00
α α α κ γ α κ	. ~ ~
where	therefore
	$v_c = 2.7$ $\alpha_x = 50$ $k/v = -0.024267$

# REDUCTIVE DECHLORINATION RATE FOR CHLORINATED ETHENES (Foc = 0.00024) MPC > MW-115 > TMP-3

# SITE OU-4 RNA TS

SHAW AFB, SOUTH CAROLINA

# The following procedures outlined by Moutoux et al. (1996)

# Step 1: Enter Observed Contaminant Concentration (mg/L)

Well	PCE	TCE	DCE	VC	Ethene	Total CAHs
MPC	90.7	718.0	5338.0	30.4	0	6177.1
MW-115	4.8	10.2	1988.0	416.0	0	2419.0
TMP-3	1.0	3.9	190.2	59.6	0	254.7

# Step 2: Compute Molar Concentrations (micromoles/Liter)

Well	M <sub>PCE</sub>	M <sub>TCE</sub>	M <sub>DCE</sub>	M <sub>VC</sub>	$M_{Ethene}$	Total M <sub>CAHs</sub>
MPC	0.55	5.47	55.06	0.49	0.00	61.56
MW-115	0.03	0.08	20.51	6.66	0.00	27.27
TMP-3	0.01	0.03	1.96	0.95	0.00	2.95

# **Step 3: Compute Carbon Equivalents**

Well	Total M <sub>CAHs</sub>	x 2	= Ceqi
MPC	61.56		123.13
MW-115	27.27		54.54
TMP-3	2.95		5.90

# Step 4: Compute Chlorine Equivalents

Well	M <sub>PCE</sub> x 4	M <sub>TCE</sub> x 3	M <sub>DCE</sub> x 2	$M_{VC}$	$S = Cleq_i$
MPC	2.19	16.40	110.13	0.49	129.20
MW-115	0.12	0.23	41.02	6.66	48.02
TMP-3	0.02	0.09	3.92	0.95	4.99

# **Step 5: Compute Corrected CAH Concentrations**

 $C_{i,corr} = C_{i-1,corr} \times (Cleq_i / Cleq_{i-1}) \times (Ceq_{i-1} / Ceq_i)$ 

Well	C <sub>i-1,corr</sub>	Cleq <sub>i</sub> / Cleq <sub>i-1</sub>	Ceq <sub>i-1</sub> / Ceq <sub>i</sub>	$C_{i,corr}$
MPC	6177.10			6177.10
MW-115	6177.10	0.37	2.26	5182.95
TMP-3	5182.95	0.10	9.24	4977.19

## Step 6: Plot Exponential Trendline of Contaminant Concentration vs. Time

	Distance From	CAH avg. velocity	CAH Travel	
Well	Source (ft)	(ft/day)*	Time (day)	$C_{i,corr}$
MPC	0	2.970	0	6177.10
MW-115	90 ·	2.970	30	5182.95
TMP-3	230	2.970	77	4977.19

<sup>\*</sup>Assumes an unretarded velocity of 3.3 feet/day below the upland terrace and a retardation coefficient of 1.11.

## **Reductive Dechlorination Rate**

 $C = C_0 e^{-kt}$ 

where:

C = Corrected Contaminant Concentration (mg/L) at time t (days)

C<sub>0</sub> = Initial Contaminant Concentration (mg/L)

k = Reductive Dechlorination Rate (days<sup>-1</sup>)

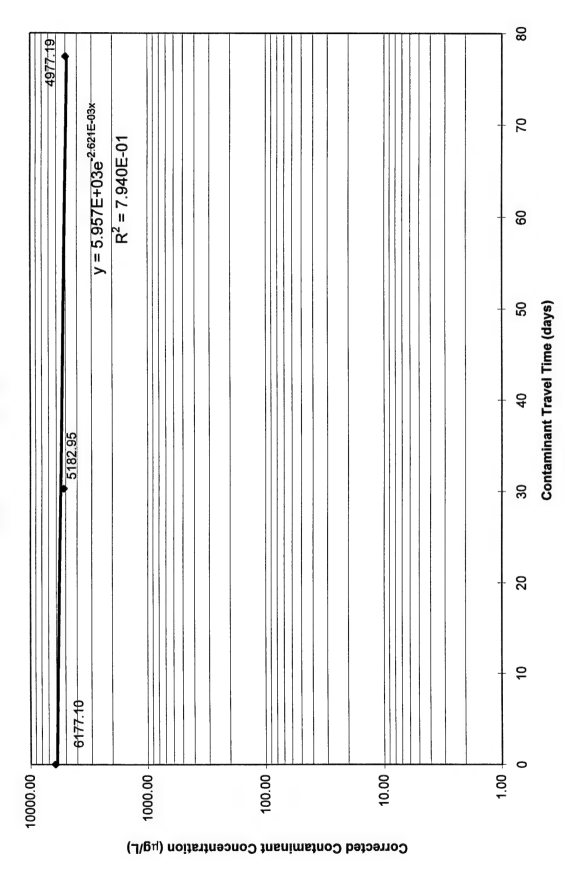
from plot:

 $y = 5957e^{-2.621E-03x}$ 

 $k = -2.621 E-03 days^{-1}$ 

REDUCTIVE DECHLORINATION RATE FOR CHLORINATED ETHENES (Foc = 0.00024) MPC > MW-115 > TMP-3

MAY, 1994 SITE OU4 RNA TS SHAW AFB, SOUTH CAROLINA



### REDUCTIVE DECHLORINATION RATE FOR CHLORINATED ETHANES (Foc = 0.00024)

### MPC > MW-115 > TMP3

SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA

### The following procedures outlined by Moutoux et al. (1996)

### Step 1: Enter Observed Contaminant Concentration (µg/L)

	Well	1,1,1-TCA	1,1-DCA	1,2-DCA	CA	Total CAHs
•	MPC	23000.0	1060.0	0.0	0.0	24060.0
	MW-115	2030.0	8700.0	0.0	0.0	10730.0
	TMP-3	60.5	1153.0	2.1	0.0	1215.6

### Step 2: Compute Molar Concentrations (micromoles/Liter)

Well	M <sub>1,1,1-TCA</sub>	M <sub>1.1-DCA</sub>	M <sub>1,2-DCA</sub>	$M_{CA}$	Total M <sub>CAHs</sub>
MPC	172.41	10.71	0.00	0.00	183.13
MW-115	15.22	87.91	0.00	0.00	103.13
TMP-3	0.45	11.65	0.02	0.00	12.13

### Step 3: Compute Carbon Equivalents

Well	Total M <sub>CAHs</sub>	x 2	= Ceqi
MPC	183.13		366.25
MW-115	103.13		206.26
TMP-3	12.13		24.25

### Step 4: Compute Chlorine Equivalents

Well	$M_{1,1,1-TCA} \times 3$	$M_{1,1-DCA} \times 2$	$M_{1,2\text{-DCA}} \times 2$	$M_{CA}$	$\Sigma = Cleq_i$
MPC	517.24	21.42	0.00	0.00	538.66
MW-115	45.65	175.83	0.00	0.00	221.48
TMP-3	1.36	23.30	0.04	0.00	24.71

### **Step 5: Compute Corrected CAH Concentrations**

 $C_{i,corr} = C_{i-1,corr} \times (Cleq_i / Cleq_{i-1}) \times (Ceq_{i-1} / Ceq_i)$ 

Well	C <sub>i-1,corr</sub>	Cleq <sub>i</sub> / Cleq <sub>i-1</sub>	Ceq <sub>i-1</sub> / Ceq <sub>i</sub>	Сі,соп
MPC	24060.00			24060.00
MW-115	24060.00	0.41	1.78	17565.86
TMP-3	17565.86	0.11	8.51	16664.88

### Step 6: Plot Exponential Trendline of Contaminant Concentration vs. Time

	Distance From	CAH avg. velocity	CAH Travel	
Well	Source (ft)	(ft/day)*	Time (day)	$C_{i,corr}$
MPC	0	2.920	0	24060.00
MW-115	30	2.920	10	17565.86
TMP-3	230	2.920	79	16664.88

<sup>\*</sup>Assumes an unretarded velocity of 3.3 feet/day and an average retardation coefficient of 1.13.

### **Reductive Dechlorination Rate**

 $C = C_0 e^{-kt}$  where:

C = Corrected Contaminant Concentration ( $\mu$ g/L) at time t (days)

 $C_0$  = Initial Contaminant Concentration ( $\mu$ g/L)

k = Reductive Dechlorination Rate (days<sup>-1</sup>)

from plot: y =

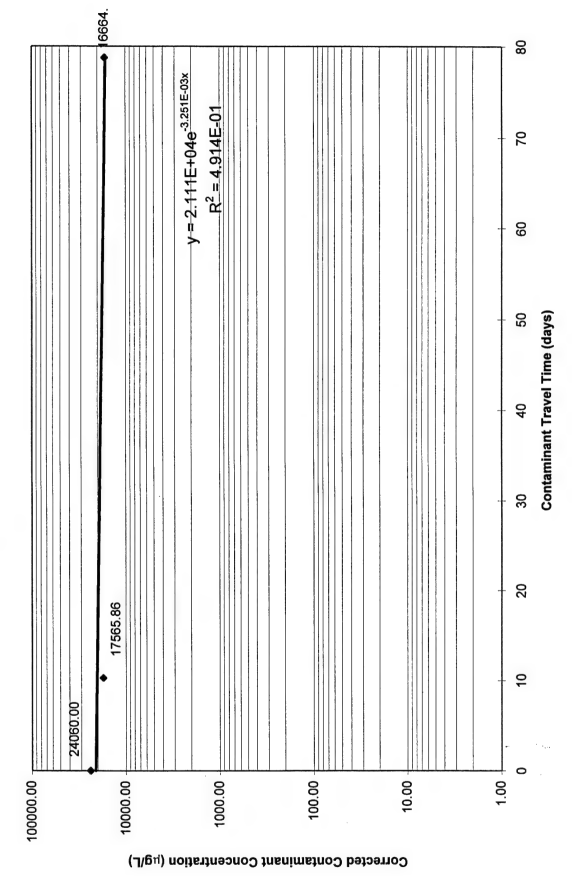
 $y = 21110e^{-3.251E-03x}$ 

 $k = -3.251E-03 days^{-1}$ 

L:\45037\OUR\TABLES\CORRATE1.XLS

## REDUCTIVE DECHLORINATION RATE FOR CHLORINATED ETHANES (Foc = 0.00024) MPC > MW-115 > TMP-3

MAY, 1997 SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA



## REDUCTIVE DECHLORINATION RATE FOR CHLORINATED BENZENES (Foc = 0.00024) MPC > MW-115 > TMP-3 SITE OU-4 RNA TS

### SHAW AFB, SOUTH CAROLINA

### The following procedures outlined by Moutoux et al. (1996)

### Step 1: Enter Observed Contaminant Concentration (µg/L)

Well	1,2-DCB	1,3-DCB	1,4-DCB	CB	Total CB's
MPC	117.0	14.1	68.2	9.2	208.5
MW-115	88.9	11.6	46.4	4.9	151.8
TMP-3	6.3	0.0	3.3	0.0	9.6

### Step 2: Compute Molar Concentrations (micromoles/Liter)

Well	M <sub>1,2-DCB</sub>	M <sub>1.3-DCB</sub>	M <sub>1,4-DCB</sub>	M <sub>CB</sub>	Total M <sub>CBs</sub>
MPC	0.80	0.10	0.46	0.08	1.44
MW-115	0.60	0.08	0.32	0.04	1.04
TMP-3	0.04	0.00	0.02	0.00	0.07

### **Step 3: Compute Carbon Equivalents**

Well	Total M <sub>CBs</sub>	x 6	= Ceqi
MPC	1.44		8.63
MW-115	1.04		6.26
TMP-3	0.07		0.39

### Step 4: Compute Chlorine Equivalents

Well	$M_{1,2\text{-DCB}} \times 2$	M <sub>1,3-DCB</sub> x 2	M <sub>1,3-DCB</sub> x 2	M <sub>CB</sub>	$\Sigma = Cleq_i$
MPC	1.59	0.19	0.93	0.08	2.79
MW-115	1.21	0.16	0.63	0.04	2.04
TMP-3	0.09	0.00	0.04	0.00	0.13

### **Step 5: Compute Corrected CAH Concentrations**

 $C_{i,corr} = C_{i-1,corr} \times (Cleq_i / Cleq_{i-1}) \times (Ceq_{i-1} / Ceq_i)$ 

	Well	C <sub>i-1,corr</sub>	Cleq <sub>i</sub> / Cleq <sub>i-1</sub>	Ceq <sub>i-1</sub> / Ceq <sub>i</sub>	$C_{i,corr}$	
•	MPC	208.50			208.50	
	MW-115	208.50	0.73	1.38	210.12	
	TMP-3	210.12	0.06	15.97	214.60	

### Step 6: Plot Exponential Trendline of Contaminant Concentration vs. Time

	Distance From	CAH avg. velocity	CAH Travel	
Well	Source (ft)	(ft/day)*	Time (day)	$C_{i,corr}$
MPC	0	2.560	0	208.50
MW-115	30	2.560	12	210.12
TMP-3	230	2.560	90	214.60

<sup>\*</sup>Assumes an unretarded velocity of 3.3 feet/day and an average retardation coefficient of 1.285.

### **Reductive Dechlorination Rate**

 $C = C_0 e^{-kt}$  where:

C = Corrected Contaminant Concentration ( $\mu$ g/L) at time t (days)

 $C_0$  = Initial Contaminant Concentration ( $\mu$ g/L)

k = Reductive Dechlorination Rate (days<sup>-1</sup>)

from plot:

 $y = 208.9e^{+3.025E-04x}$ 

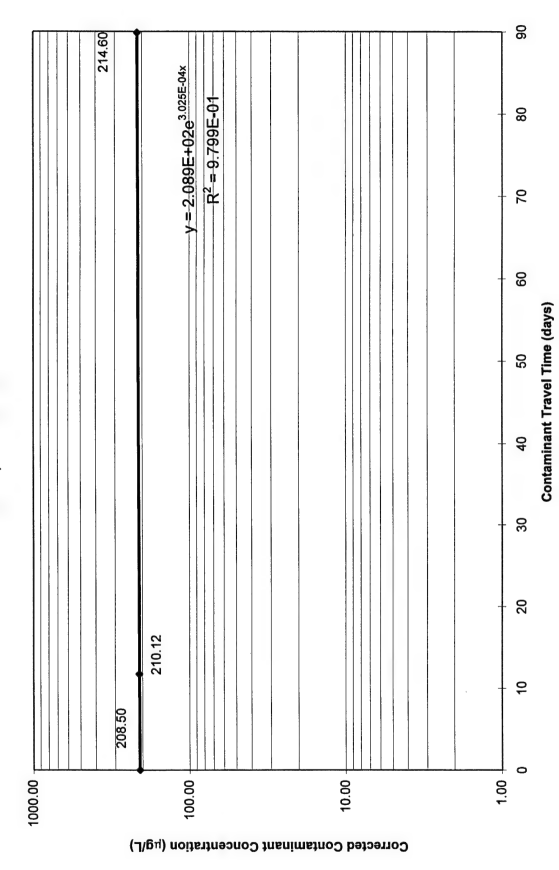
 $k = +3.025E-04 days^{-1}$ 

(ERRONEOUS RATE - INCREASING IN VALUE)

L:\45037\OU4\TABLES\CORRATE1.XLS

# REDUCTIVE DECHLORINATION RATE FOR DICHLOROBENZENES TO MONOCHLOROBENZENE (Foc = 0.00024) MPC > MW-115 > TMP-3

MAY, 1997 SITE OU-4 RNA TS SHAW AFB, SOUTH CAROLINA



### APPENDIX D

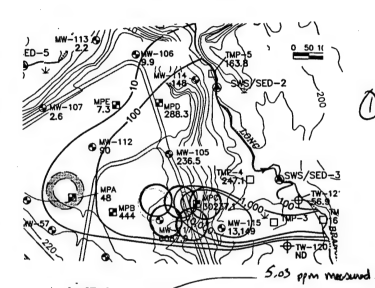
### MODEL INPUT FILES

### APPENDIX E

## COST CALCULATIONS PERTINENT TO REMEDIAL ALTERNATIVES SELECTION

Estimating CAH sail goo cone. i polardial SNE System size in some area near MPC/MW-117.

According to the pilot scale chemical oxidesteni lest (IT, 1997), a combined soil gos ancentration of 5,830 ppb, or 5.03 ppm was measured at mw-Pl. This location was approximately 30' Alw of MW-117. This was prior to chemical exidetion treatment. Since



200 FEET

TOTAL VOCS PRIOR TO TREATMENT

SHAW AIR FORCE BASE SUMTER, SOUTH CAROLINA

Assume that a 50' Radius . I Influence can be achieved. Assume we went to have at least 10 pore volume exchanges for each radius at influence

Design SVE Blaver Size !

Estinto VOC'S Removed / Day

what is the volume in liters for 10 pore volumes per well per dany

V= 10' x 10' x 0.3 x 11 (50)

V= 58,905#13 x 28.316 = 1.649 x 1061

(2) what is the total values for all wellc?

5 x 1.649 x 106L = 8.2467 x 106L

3) If we ossume lopping initial voc core, , what will we initially resover pardey?

8.2467x101 x 10m3 = 8.2467x107m

= 82 kg/deg.

This seems high, bout may to exhibites in the mitil few unters.

(4) what blown rate myst we need; or size?

@ 8.2467 x 10° L x 0.03532H3 = 291,273 A3/dey = = 202.3 sefm We a moder with Double apaidy - ROTRON DR8, max of 400 scfm. MERCED 8 ACT

### LTM 1 Cost Estimate Shaw AFB, Site OU4 722450.3705

Author: RTH Date: 2/17/98 Checked by:

Date:

### **Summary of Capital and Present Worth Costs**

### **Capital Costs**

Design/Construct 4 LTM Wells and initiate LTM Program in 1998

P/F i=7% n=0

**Monitoring Costs** 

Annual Monitoring of 11 wells and 6 surface water stations, 1998-2003

(5 events)

Cost per Event	\$22,170	
P/A i=7%, n=0	1998	\$22,170
P/A i=7%, n=1	1999	\$20,720
P/A i=7%, n=2	2000	\$19,364
P/A i=7%, n=3	2001	\$18,097
P/A i=7%, n=4	2002	\$16,913

**Total Present Worth Cost** 

\$97,264

\$12,449

Biennial Monitoring of 11 wells and 6 surface water stations, 2004-2018 (8 events)

Cost per Event	\$22,170	
P/A i=7%, n=6	2004	\$14,773
P/A i=7%, n=8	2006	\$12,903
P/A i=7%, n=10	2008	\$11,270
P/A i=7%, n=12	2010	\$9,844
P/A i=7%, n=14	2012	\$8,598
P/A i=7%, n=16	2014	\$7,510
P/A i=7%, n=18	2016	\$6,559
P/A i=7%, n=20	2018	\$5,729

**Total Present Worth Cost** 

\$57,388

Site Management every year (20 years)

**Annual Cost** 

\$6,000

P/A i=7% n=20

PWF = 10.8355273

Present Worth Cost

\$65,013

**Total Capital and Present Worth Costs of LTM Program** 

\$232,114

### LTM 1 Cost Estimate SITE OU4 RNA TS 722450.37050

Author: RTH Date: 2/17/98 Checked by:

Date:

GW and Surface Water Sampling - 11 wells/6 surface water stations sampled annually till 2002 11 wells/6 surface water stations sampled biennually till 2018.

Sampling Labor 11 Long-Term Monitoring Wells 6 Surface Water Sample 4 QA/QC 21 Total Samples	90 hours	x	\$60	/hour	\$5,400
Analytical Subcontractor					
	21 CAHs	•	\$150	/each	\$3,150
	13 Metha	ane/Ethene	\$100	/each	\$1,300
	13 Field	Parameters	\$20	/each	\$260
Supplies			\$700	lump sum	\$700
Travel			\$1,300	lump sum	\$1,300
Data Management (40 hr x \$60hr)			\$2,400		\$2,400
Data Validation (20 hr x \$60/hr)			\$1,200		\$1,200
Reporting/Project Management Labor					
Word Processing	10	hours x	\$25	/hour	\$250
CADD	15	hours x	\$50	/hour	\$750
Reproduction	8	hours x	\$20	/hour	\$160
Staff Level	50	hours x	•	/hour	\$3,000
Proj. Manager	20	hours x	-	/hour	\$1,600
Editor	5	hours x	\$60	/hour	\$300
Reporting/Project Management ODCs			\$400	lump sum	\$400

\$22,170

Total for 1 Sampling Event

## Design/Construct 4 LTM Wells & Initiate LTM Program (4 shallow wells installed)

Shaw AFB Site OU-4

Standard Rate Schedule

Standard Rate Schedu	пС							
Billing		Billing		Install New		Subcon-		
Category	(4		Task 1	LTM/POC	Task 2	tracting	Task 3	Reporting
Cost Code/(Billing Ca	ategory)	Rate	(hrs)	Wells (\$)	(hrs)		(hrs)	& PM (\$)
Word Processor 88	/(15)	\$30	0	\$0	5	\$150	5	\$150
CADD Operator 58	3/(25)	\$47	0	\$0	0	\$0	0	\$0
	2/(50)	\$40	8	\$320	0	\$0	0	\$0
Staff Level 16	/(65)	\$57	20	\$1,140	0	\$0	0	\$0
Project Level 12	/(70)	\$65	8	\$520	0	\$0	0	\$0
	/(80)	\$85	2	\$170	0	\$0	0	\$0
Principal 02	2/(85)	\$97	0	\$0	0	\$0	0	\$0
Total Labor (hrs  \$)			38	\$2,150	0	\$0	0	\$0
ODCs								
Phone				\$20		\$0		\$0
Photocopy				\$20 \$10		\$0 \$0		\$0 \$0
Mail				\$0		\$0		\$0 \$0
Computer				\$0		\$0		\$0
CAD				\$0		\$0		\$0
WP `				\$0		\$0		\$0
Travel				\$600		\$0		\$0
Per Diem				\$100		\$0		\$0
Eqpt. & Supplies				\$700		\$0		\$0
Total ODCs				\$1,430		\$0		\$0
Outside Services								
LTM/POC Well Insta	allation Costs	•		\$8,269		\$0		\$0
Surveying	andion Costs	,		\$600		\$0		\$0 \$0
Other: Maintain Inst	itutional Con	trols		\$0		\$0		\$0
Total Outside Service	es			\$8,869		\$0		\$0

Proposal Estimate	Task 1	Task 2	Task 3
Labor	\$2,150	\$0	\$0
ODC's	\$1,430	\$0	\$0
Outside Services	\$8,869	\$0	\$0
Total by Task	\$12,449	\$0	\$0
Total Labor	\$2,150		
Total ODCs	\$1,430		
Total Outside Services	\$8,869		
Total Project	\$12,449		

Task 1: Install New LTM wells (4 shallow)

Task 2: Subcontracting/Permitting

Task 3: Reporting/PM per Event.

## Shaw AFB, Site OU-4 Backup Calculations

1

Long-Term Monitoring Well Installation	allation						
		Cost calculations					
Misc calculations		Description	Unit	Qty.	Qty.   Unit Price   Subtotal	Subtotal	Total
Number of LTM wells:		Well Installation					\$ 8,269
Number of wells:	2	Mobilization	ea	-	\$ 500	\$ 500	
Depth each:	35 ft	Well Installation	ln ft	120	\$	\$ 7,200	
		Soil Disposal	drum	9	\$ 100	\$ 269	
Number of wells:	2						
Depth each:	25 ft						
Total Depth:	120						

Installation of SVE Wells								
		Cost calculations						
Misc calculations		Description	Unit	Qty.	Qty. Unit Price	Subtotal	_	Total
Number of LTM wells:		Well Installation					€9	5,356
Number of wells:	5	Mobilization	ea	-	\$ 500 \$	\$ 500		
Depth each:	15 ft	Well Installation	ln ft	75	<del>69</del>	60 \$ 4,500		
Total Depth:	75	Soil Disposal	drum	4	\$ 100	100 \$ 356		

### Remedial Alternative 2 Cost Estimate SHAW AFB, SOUTH CAROLINA 722450.37050

Author: RTH Date: 2/17/98 Checked by:

Date:

<b>Summary of Present Worth Capital Costs</b>	(assuming system installation in 1998)	

Installation of Vapor Extraction and Treatment System

\$115,856 P/A i=7%, n=0

\$115,856

Total

\$115.856

### Summary of Present Worth Annual + Biennial + Future Costs (assuming 5-yr operating period 1998-2003)

O&M for SVE system (5 years)

Annual Cost \$55,520 1998 n=0 \$55,520 1999 n=1 \$51,888 2000 n=2 \$48,493 2001 n=3 \$45,321 2002 n=4 \$42,356

Future Costs (replace regenerative motor once in 2001)

cost per event =

\$2,500

2001 n=3

\$2,041

Total =

\$2,041

**Total Present Worth Cost** 

\$245,619

Total Present Worth Cost of LTM Program (from Alternative 1)

(includes site management)

\$232,114

**Total Annual and Biennial Costs** 

\$477,733

Total Capital + Annual + Biennial + Future Costs for Alternative 2

\$593,589

### Remedial Alternative 2 Cost Estimate SHAW AFB, SOUTH CAROLINA 722450.37050

Author: RTH Date: 2/17/98 Checked by:

Date:

Capital Costs						
Installation of the	Vapor Extraction System SVE Pilot Test (assume existing wells used)				lump sum	\$25,000
	Design/Procure System	150	hours x	\$70	/hour	\$10,500
	Install SVE Wells			\$5,356	lump sum	\$5,356
	Blower/Shed/Accessories			\$20,000	lump sum	\$20,000
	Thermal Oxidizer Vapor Treatment System (Includes Nat. Gas Costs	_	months x	\$5,500	/month	\$16,500
	Carbon Adsorption Vapor Treatment System			\$12,000	lump sum	\$12,000
	Electrical Subcontractor			\$5,000	lump sum	\$5,000
	Per Diem	20	days x	\$145	/day	\$2,900
	Travel (1 round trip)			\$1,000	lump sum	\$1,000
	O & M Manual Preparation	40	hours x	\$65	/hour	\$2,600
	Emissions Permitting			\$2,000	lump sum	\$2,000
	Installation/startup Labor	200	hours x	\$65	/hour	\$13,000
					Subtotal	\$115,856

### Remedial Alternative 2 Cost Estimate SHAW AFB, SOUTH CAROLINA 722450.37050

Author: RTH Date: 2/17/98 Checked by:

Date:

### Annual Operating Costs--SVE system (5 years)

Labor	
Weekly System Checks (2 hr x 52 wk/yr x \$60/hr)	\$6,240
Monthly Sampling (12 hr x 12 mo x \$60/hr)	\$8,640
Monthly Reporting (8 hr x 12 mo x \$60/hr)	\$5,760
Maintenance (8 hr x 12 mo x \$60/hr)	\$5,760
Analytical	
VOCs (air) (2/mo x 12 mo x \$250/ea)	\$6,000
Sample ship. (2/mo x 12 mo x \$30/ea)	\$720
Supplies	
Install and Dispose of GAC (assume 2 replacements of 2,000 lb. of GAC/year)	\$9,400
misc. (lump sum)	\$1,000
Travel (52 trips from Raleigh, NC per year @ \$100/ea)	\$5,200
Electricity	\$5,000
Site Management (Included in LTM portion of Alternative 1)	\$0
Recovered Solvent Disposal from Knock Out Tanks (\$150/mo x 12 mo)	\$1,800
subtotal	\$55,520

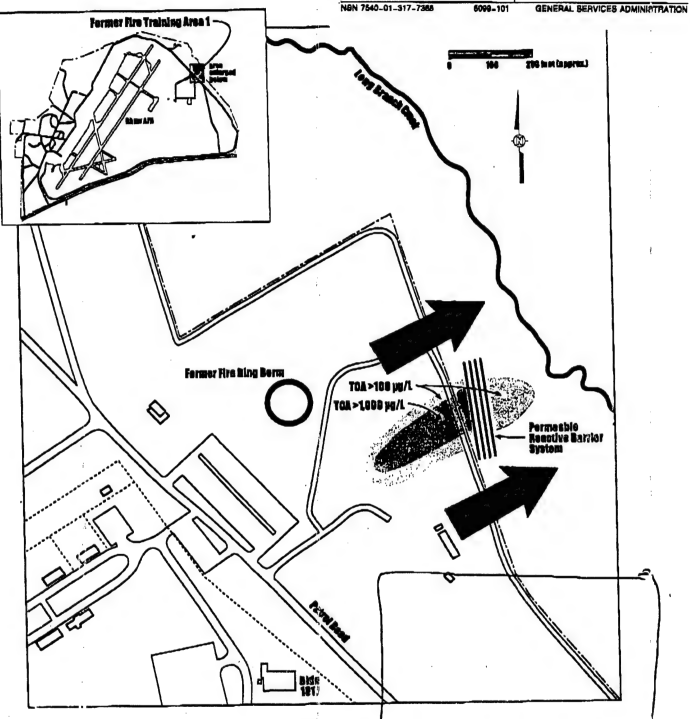
### APPENDIX E

## APPROXIMATE LOCATION OF IRON FILINGS WALL (FORWARDED BY MR. RICK ROLLER, 20 CES/CEV ON 9/10/99)

FAX TRANSMITTAL

NON 7540-01-317-7368

6099-101



## APPENDIX F REPORT COMMENTS AND RESPONSE TO COMMENTS

### **PARSONS**

Parsons Engineering Science, Inc. • A Unit of Parsons Infrastructure & Technology Group Inc.

1700 Broadway, Suite 900 • Denver, Colorado 80290 • (303) 831-8100 • Fax: (303) 831-8208 • www.parsons.com

July 7, 1999

Mr. Jerry Hansen Technical Program Manager AFCEE/ERT 3207 North Road, Bldg. 532 Brooks AFB, TX 78235-5363 (210) 536-4353

Subject: Response to Comments for the Draft Remediation by Natural Attenuation Treatability Study for OU-4 (Site FT-1), Shaw Air Force Base, South Carolina.

### Dear Mr. Hansen:

Mr. Jon Atkinson of HQ AFCEE/ERC at Brooks AFB, Texas submitted formal comments for the subject report. The following is a summary of the response to comments provided by Parsons ES:

## COMMENTS BY: MR. JON ATKINSON, HQ AFCEE/ERC (25 FEBRUARY 1999)

**Comment 1:** Page V: Suggest listing the appendices in the table of contents.

**Response 1:** Agreed. The appendices will be listed in the table of contents.

Comment 2: Page 1-2, Sect. 1.1, Sent. 1: Suggest expanding to add "and technical feasibility" after "defensibility".

Response 2: Agreed. The referenced sentence now reads, "...to conduct site characterization and groundwater modeling to evaluate the scientific defensibility and technical feasibility of RNA with long-term monitoring (LTM)..."

Comment 3: Page 2-1, Sec. 2, Bullet 3, Line 5: Recommend changing "aromatic acids" to "aromatic hydrocarbons".

Response 3: The reference to aromatic acids was misplaced and misreferenced. The sentence now reads, "...sulfate, total organic carbon (TOC), dissolved hydrogen, and methane/ethane/ethene...."

Comment 4: Page 3-1, Sec. 3.1, Par. 2, Sent. 2: Suggest stating whether Long Branch Creek is perennial or ephemeral.

- **Response 4:** Agreed. The sentence now reads: "The small perennial stream, Long Branch Creek, and an unnamed southwest drainage ditch both border the Base..."
- Comment 5: Pages 3-7 and 3-8, Figs 3.3 and 3.4: Suggest adding the vertical exaggeration specifications for the cross sections.
- **Response 5:** Agreed. References for the vertical exaggeration in Figures 3.3 and 3.4 were added.
- Comment 6: Page 3-13, Sec. 3.3.2.2, Sent. 1: Typically pumping tests are performed for periods ranging from 24 to 72 hours. Consequently, should "26-minute" be "26-hour"?
- **Response 6:** The reference to a "26-minute" pumping test is correct as stated. This fact was confirmed with Mr. Rick Roller of Shaw AFB on 5/21/99.
- Comment 7: Page 4-6, sec. 4.1.2, Sent. 2: To correct terminology, recommend changing "groundwater table" to "water table."
- **Response 7:** Agreed. "Groundwater table" was changed to "water table" to maintain consistency of the use of this term in Section 4.1.2.
- Comment 8: Page 4-23, Sec. 4.3.1, Par. 1, Last Line: To better reflect accuracy and certainty of analytical methods here and throughout Section 4.3, suggest reporting results to no more than three significant figures (1930 μg/L).
- Response 8: It is the laboratory's responsibility to determine the significant figure placement for analytical data provided in hardcopy format. The data hardcopy is the legally defensible form of data provided to Parsons ES by the laboratory and Parsons ES has chosen to maintain consistency throughout the report by using hardcopy values in all text and tables. Furthermore, utilization of nonrounded hardcopy results helps to avoid reader confusion when reviewing tables and text.
- Page 4-24, Table 4.3: To better reflect accuracy and certainty of analytical methods, suggest reporting results to no more than three significant figures (round 340.6 to 341. μg/L.
- **Response 9:** See Response 8.
- Comment 10: Page 4-35, Table 4.5: To better reflect accuracy and certainty of analytical methods, suggest reporting results to no more than three significant figures (round 340.6 to 341  $\mu$ g/L)
- **Response 10:** See Response 8.

- Comment 11: Pages 4-50 and 4-53, Sec. 4.4.2.1, Line 10: To improve sentence structure, recommend inserting "from" after "resulted."
- Response 11: Agreed. The change was made and the next now reads, "...contaminant plume (MPC and MW-117) may have resulted *from* the chemical oxidation pilot test, which caused a large release...."
- Comment 12: Page 4-53 and 4-54, Sec. 4.4.2.1, Line 14: "19.02 mg/L" should be "19.02 nM/L."
- **Response 12:** Agreed. The referenced text now reads "A dissolved hydrogen concentration of 19.02 *nM/L* was measured at MW-112A which is..."
- Comment 13: Page 4-61, Sec. 4.6, Par. 3, Line 1: "coocurrence" probably should be "occurrence."
- Response 13: Agreed. The sentence now reads, "The occurrence of CB and DCB contamination indicates...."
- Comment 14: Pages 6-29 and 6-36, Tables 6.3 and 6.4: Four wells are cited under capital costs; however, Section 7.2.1 (p. 7-2) state that three new LTM wells are recommended. This inconsistency needs to be corrected.
- Response 14: Parsons ES proposed four new wells at OU4. These four proposed wells include three LTM wells and one sentry well. The proposed sentry well location was accidentally omitted from Figure 7.1 and will be replaced in the final document. Tables 6.3 and 6.4 were clarified by the modification of the first line of descriptive text in each table from, "Design/construct four LTM wells in 1998," to, "Design/construct three LTM wells and one sentry well in 1998." Furthermore, the costing section for Alternative 1 (Section 6.4.1.3) was amended to include a reference to the construction of a sentry well.
- Comment 15: Page 7-2, Sec. 7.2, Line 5: To correct the grammar, "consist" should be changed to "consists."
- **Response 15:** Agreed. This change was made to the Final TS.
- Comment 16: Page 7-2, Sec. 7.2.1: MW-112 is listed as a proposed LTM well; however, it is not depicted as a proposed LTM well on Figure 7.1.
- **Response 16:** Thank you. Monitoring well MW-112 was marked as a proposed LTM well in Figure 7.1.
- Comment 17: Page 7-4, Sec. 7.2.2, Sent. 2: Although this sentence states that a sentry well is proposed about 300 feet east of Long Branch Creek, this sentry

well is not depicted on Figure 7.1. This inconsistency should be resolved.

**Response 17:** Agreed. See Response 14.

Comment 18: Page 8-2, Sec. 8, Par. 1, Line 5: Suggest changing "DHEC (1997)" to "SCDHEC (1997)."

**Response 18:** Agreed. DHEC was changed to SCDHEC in the above reference and in other similar references in the TS report.

Comment 19: Page 8-4, Sec. 8

a. Par. 1, Line 1: "affects" should be "effects."

b. Par. 2, Sent. 1: The recommendation of four proposed new monitoring wells contradicts Section 7.2.1 and Figure 7.1 which contain three proposed LTM wells. This inconsistency needs to be resolved.

### Response 19:

a. Agreed. "affects" was changed to "effects."

b. This reference is correct. Please see Response 14.

Comment 20: Page 9-6, Sec. 9, Ref. 5: Suggest inserting "South Carolina" in front of "Department."

**Response 20:** Agreed. This change was made.

### COMMENTS BY: MR. RICHARD C. ROLLER, 20 CES/CEV (18 MARCH 1999)

Comment 1: Comments are provided to accept the referenced document as final by Shaw AFB, SC. The statement at the top of page ES-3 should reflect that AFCEE as a result of this investigation and study recommends this particular solution. It should not bind the Air Force, inclusive of Shaw AFB.

Response 1: Agreed. The referenced statement now reads ". The Air Force Center for Environmental Excellence (AFCEE) therefore recommends implementation of soil vapor extraction (SVE)..."

Comment a: Shaw AFB addressed cleanup of chlorinated solvents at the OU-4 site by construction of a Permeable Reactive Barrier Wall in November 1998. It is requested that this information be referenced in the executive

summary and that the conclusions and recommendation on page ES-3 should be reflective of the current decisions at the site.

### Response a:

Agreed. Reference to the installation of a permeable reactive barrier wall will be made in the executive summary and in Section 6. A brief analysis of the potential benefits and disadvantages of a permeable reaction wall was given in Section 6.2.4 of the TS report. The conceptual purpose of a reactive wall is to degrade contaminants before they can enter into Long Branch Creek. A reactive barrier wall is not intended to treat the source LNAPL at the site. Therefore, groundwater contamination may persist for decades, although mitigated by the wall before entry into Long Branch Creek. AFCEE continues to recommend source delineation and treatment and RNA with LTM as the preferred remedial alternative.

### Comment b:

The present status of the site is to monitor the effectiveness of the reactive wall and to allow natural attenuation and phytoremediation to address the contamination downgradient of the wall. The base is conducting additional sampling to evaluate the current state of DRO contamination and also to make a decision on the deeper chlorinated solvent plume that could potentially go under the reactive wall.

### Response b:

Reference to the present status will be made in the executive summary and in Section 6.

### Comment c:

Other cleanup systems may be incorporated as result of this investigation. The AFCEE study might reference this work and potential impact on the recommendations, although it is understood that the AFCEE study was accomplished for a specific purpose.

### Response c:

Without knowledge of the specific cleanup systems being considered by Shaw AFB, Parsons ES cannot assess potential impacts to the remediation period and LTM program. Conversations with Mr. Rick Roller, 20 CES/CEV, Shaw AFB indicates that potential future remediation systems would apply to the diesel range organic (DRO) plume of the northeast/east plume lobe. The selection of a treatment technology for the DRO plume will not be determined until after an additional site investigation is performed. Groundwater modeling would be the most useful tool for assessing the impacts of additional remedial systems on the DRO plume once additional groundwater data is gathered. However, additional groundwater modeling is beyond the scope and schedule of the current project. Parsons ES recommends that the impacts of additional remedial systems be evaluated in other reports.

Mr. Jerry Hansen July 7, 1999 Page 6

If you have any questions or comments regarding the response to comments, please do not hesitate to contact Mr. Bruce Henry or Mr. Todd Herrington at (303) 831-8100.

Sincerely,

PARSONS ENGINEERING SCIENCE, INC.

Bruce M. Henry Project Manager

**Enclosures**